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(54) Title: ANTISENSE OLIGONUCLEOTIDES AS ANTIBACTERIAL AGENTS

(57) Abstract

A novel method is provided that teaches the therapeutic use of nuclease resistant oligonucleotides for treating animals having an infection caused by a pathogenic bacterium. The method involves the integration of (1) methods for selecting the correct oligonucleotide, (2) synthesis and purification of nuclease resistant oligonucleotides, and (3) methods for in vitro analysis of potential antimicrobial oligonucleotides. The described oligonucleotides may comprise modified backbones, sugar residues, bases, or mixtures and have been subject to purification resulting in oligonucleotides that are capable of inhibiting the growth of a broad spectrum of clinically relevant bacterial species.

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ANTISENSE OLIGONUCLEOTIDES AS ANTIBACTERIAL AGENTS

The present application claims priority to United States Patent Application Serial No. 08/685,575, filed July 24, 1996.

5

FIELD OF THE INVENTION

The present invention is directed to methods for treating an animal, including a human, having a bacterial infection which comprise administering an oligonucleotide

10 specifically targeted to, or otherwise capable of interacting with, a bacterial sequence, or nucleic acid binding protein. The antibacterial oligonucleotide inhibits the growth of the bacteria, blocks the expression of virulence factors or genes involved in the transfer of genetic information, or kills the

15 bacteria. Alternatively, the oligonucleotide may also be targeted to an antibiotic resistance gene in order to render the bacteria sensitive to an otherwise ineffective antibiotic. The invention also relates to nuclease resistant oligonucleotides that are effective in inhibiting the growth

20 of, or killing, pathogenic bacteria.

1.0. BACKGROUND TO THE INVENTION

1.1. Antibiotic Prior Art

Pathogenic bacteria responsible for infectious diseases

25 were once thought to be totally under control through the use
of a battery of antibiotics such as penicillin, streptomycin,
tetracycline, and others. However, since the widespread use
of antibiotics began in the 1950s, more and more bacteria
resistant to one or more antibiotics have arisen. Multiple

30 drug resistant strains are increasingly common, particularly
in hospitals.

Currently, nosocomial Staphylococcal infections exhibit multiple drug resistance. See, for example, Archer et al., Antimicrob. Agents Chemother. 38:2231-2237 (1994). At this time, the remaining antibiotic that demonstrates the ability to kill Staphylococci is vancomycin. Strains of Enterococci that are vancomycin resistant have already been isolated and

reported by Zabransky et al., J. Clin. Microbiol. 33(4):791-793 (1995). Furthermore, transfer of resistance from Enterococci to Staphylococci has been previously documented by Woodford et al., J. Antimicrob. Chemother. 35:179-184

- 5 (1995). Streptococcus pneumoniae is a leading cause of morbidity and mortality in the United States (M.M.W.R., Feb. 16, 1996, Vol. 45, No. RR-1). Each year these bacteria cause 3,000 cases of meningitis, 50,000 cases of bacteremia, 500,000 cases of pneumonia, and 7,000,000 cases of otitis
- 10 media. Case fatality rates are greater than 40% for bacteremia and greater than 55% for meningitis, despite antibiotic therapy. In the past, Streptococcus pneumoniae were uniformly susceptible to antibiotics; however, antibiotic resistant strains have emerged and are becoming 15 widespread in some communities.

In addition, there are instances where antibiotic resistance is not an issue, yet a particular bacteria remains refractory to treatment using conventional antibiotics. Such is the case with *Escherichia coli* 0157:H7, the causative

- 20 agent for food poisoning and death from undercooked meat. The Department of Agriculture estimates that 10 people die each day and another 14,000 become ill due to this bacteria. Unfortunately, conventional antibiotics are completely ineffective against this organism.
- 25 The history of antibiotic treatment of pathogenic bacteria is cyclical. Bacteria are remarkably adaptive organisms, and, for each new antibiotic that has been developed, resistant bacterial strains arise through the widespread use of the antibiotic. Thus, there is a constant
- of antibiotic resistant bacteria. Traditional methods of developing new antibiotics have slowed, and in the past two years only one new antibiotic has been approved by the FDA. Furthermore, according to Kristinsson (Microb. Drug
- 35 Resistance $\underline{1}(2):121$ (1995)), "There are no new antimicrobial classes with activity against resistant Gram positives on the horizon."

1.2. Antisense Nucleotide Art

Antisense polynucleotides are useful for specifically inhibiting unwanted gene expression in mammalian cells. They can be used to hybridize to and inhibit the function of an 5 RNA, typically a messenger RNA, by activating RNase H or physically blocking the binding of ribosomes or proteins, thus preventing translation of the mRNA. Antisense oligonucleotides also include RNAs with catalytic activity (ribozymes), which can selectively bind to complementary

10 sequences on a target RNA and physically destroy the target by mediating a cleavage reaction.

Antisense oligonucleotides that bind to the DNA at the

correct location can also prevent the DNA from being transcribed into RNA. These antigene oligonucleotides are 15 believed to bind to double-stranded DNA (forming triple-stranded DNA) and thereby inhibit gene expression.

1.3. Antisense Nucleotides For Therapy

The use of antisense oligonucleotides has emerged as a 20 powerful new approach for the treatment of certain diseases. However, the preponderance of the work to date has focused on the use of antisense oligonucleotides as antiviral agents or as anticancer agents (Wickstrom, E., Ed., Prospects for Antisense Nucleic Acid Therapy of Cancer and AIDS, New York:

25 Wiley-Liss, 1991; Crooke, S.T. and Lebleu, B., Eds.,
Antisense Research and Applications, Boca Raton: CRC Press,
1993, pp. 154-182; Baserga, R. and Denhardt, D.T., 1992,
Antisense Strategies, New York: The New York Academy of
Sciences, Vol. 660; Murray, J.A.H., Ed., Antisense RNA and
30 DNA, New York: Wiley-Liss. 1993).

There have been numerous disclosures of the use of antisense oligonucleotides as antiviral agents. For example, Agrawal et al. report phosphoramidate and phosphorothioate oligonucleotides as antisense inhibitors of HIV (Agrawal et

35 al., Proc. Natl. Acad. Sci. USA <u>85</u>:7079-7083 (1988)).

Zamecnik et al. disclose antisense oligonucleotides as inhibitors of Rous sarcoma virus replication in chicken

fibroblasts (Zamecnik et al., Proc. Natl. Acad. Sci. USA 83:4143-4146 (1986)).

There seem to be few to no toxicity problems associated with the use of antisense oligonucleotides as drugs to treat disease. To date, no dose limiting toxicities of phosphorothioate antisense oligonucleotides have been detected in man (Crooke, S.T., "Progress in Oligonucleotide Therapeutics," Abstracts American Association for Cancer Research, March 18-22, 1995; Crooke, S.T., "Progress in

- 10 Oligonucleotide Therapeutics, "Abstracts Oligonucleotide-Based Therapeutics, February 9-10, 1995), and phosphorothicate oligonucleotides have been found to have no effect on developing embryos (Guadette et al., Antisense Res. Devel. 3:391-397 (1993)). In fact, under an emergency IND
- 15 approval, a 19-year-old male received 700 mg of an antisense phosphorothicate oligonucleotide to treat acute myeloblastic leukemia (Bayever et al., Antisense Res. Devel. 2:109-110 (1992)). There were no changes in pulse, respiratory rate, blood pressure, fever, mucositis, or diarrhea in the patient.
- 20 In addition, no neurological, cardiovascular, respiratory, renal, skin or nephrourological toxicities were observed. It was concluded that systemic administration of a phosphorothicate antisense oligonucleotide to humans achieves adequate bioavailability of the drug to target tissues
- 25 without major toxicity. In a follow up study, the antisense phosphorothicate oligonucleotides were given to five patients with acute myeloblastic leukemia. After systemic intravenous administration of the oligonucleotide, no toxic effects were seen. See Fig. 1 of Bayever et al., Antisense Res. Devel.
- 30 3:383-390 (1993). The authors concluded that the favorable pharmacokinetics observed support the use of phosphorothicate oligonucleotides as potential gene specific therapeutic agents.

1.4. The Transport Problem For Oligonucleotides

While the use of antisense oligonucleotides as antiviral agents has been described (Agrawal et al., Pat. No. 5,194,428, issued March 16, 1993), no significant progress 5 has been made in the therapeutic use of antisense oligonucleotides to treat bacterial infection. In fact, at a recent meeting on Antibiotic Discovery addressing the current state of the art, there were no talks or discussions scheduled regarding the use of antisense oligonucleotides to 10 treat bacterial infections, although the use of antisense oligonucleotides as antiviral agents was scheduled for discussion ("Antibiotic Discovery," Abstracts International Business Communications, June 26-27, 1995).

Logically, the use of synthetic oligonucleotides should 15 be advantageous as an approach to treating bacterial infection because sequences can be specifically designed to inhibit bacterial growth while not interfering with the metabolism of mammalian cells.

In addition, oligonucleotides have been shown to
20 nonspecifically stimulate the immune system (Yamamoto et al.,
Antisense Res. Devel. 4:119-122 (1994); Krieg et al., Nature
374:546-549 (1995)). Since current antibiotics generally
function by arresting bacterial growth until the immune
system can respond to the infection (Myrvik, Fundamentals of

- 25 Medical Bacteriology, 1974, Lea & Febiger, Publishers), the use of oligonucleotides as antibiotics may provide both a nonspecific stimulation of the immune system as well as the relatively specific inhibition of the growth of a particular bacteria.
- Furthermore, infectious bacteria have been shown to become sequestered in the liver and spleen in clinical infections (Wilson, G.S. and Miles, A.A., Eds., Topley and Wilson's Principles of Bacteriology and Immunology, Williams & Wilkins, Publishers, 1964). Oligonucleotides, or more
- 35 specifically S-oligonucleotides (phosphorothicate substituted), have also been shown to accumulate in these organs (Agrawal et al., Proc. Natl. Acad. Sci. USA 88:7595-

7599 (1991)). Therefore, the use of antisense oligonucleotides should be ideally suited to the treatment of bacterial infections involving the liver and spleen as well as systemic bacteremia and septicemia.

- The rigid cellular architecture of the prokaryote has been viewed as a barrier to oligonucleotide uptake by bacterial cells (Chrisey et al., Antisense Res. Devel. 3:367-381 (1993)). In fact, reports of antisense oligonucleotidemediated gene inhibition in bacteria have attempted to
- 10 circumvent the perceived problem of the rigid cell wall by conducting experiments in cell-wall deficient strains (Jayaraman et al., Proc. Natl. Acad. Sci. USA 78:1537-1541 (1981)), in competent bacterial cells (Ciferri et al., J. Bacteriol. 104:684-688 (1970)), in heat-shock permeabilized
- 15 bacteria (Gasparro et al., Antisense Res. Devel. 1:117-140 (1991)), in hypertonic solutions (Chrisey et al., Antisense Res. Devel. 3:367-381 (1993)), and using PEG-modified oligonucleotides (Rahman et al., Antisense Res. Devel. 1:319-327 (1991)), none of which has relevance to treating clinical bacterial infections.

Lupski et al., Pat. No. 5,294,533 ('533 patent), stated that antisense oligonucleotides can preferentially inhibit the growth of Gram negative and Gram positive bacteria in a mixed culture of Gram negative and Gram positive bacteria.

- 25 Lupski et al. also taught that end-capped oligonucleotides should be used (see column 4, lines 39-42), but since end-capping does not provide protection from intracellular endonucleases (see the discussion of Hoke et al. above), one skilled in the art would not expect the method of Lupski et
- 30 al. to work. Thus, the '533 patent does not provide an enabling description of the use of antisense oligonucleotides to inhibit the growth of bacteria in vivo in mammals.

Moreover, the '533 patent did not disclose the genotype of the bacteria used in the study. Thus, there is no way to 35 establish whether clinical isolates were used or permeability enhanced bacterial mutants were used. Additionally, the '533 patent does not provide adequate teaching to allow one to

discern whether or not the described bacteria had been previously rendered competent by established prior art methods. In view of this lack of disclosure, the '533 patent does not teach methods that are broadly applicable to 5 clinically significant bacterial infections in mammals.

The prior art teaches the inherent difficulty of successfully using oligonucleotides to inhibit the growth of intact bacteria (Jayaraman et al. and Ciferri et al.), and the '533 patent does not provide sufficient disclosure to

- 10 refute the clear teaching in the prior art. Instead, the '533 patent simply states that: "A small 10-29 mer antisense oligonucleotide that is delivered to a bacteria is rapidly transported into the bacterial cells." This statement is clearly contrary to what is taught by the prior art.
- The prior art has never conclusively established that the growth of wild type bacteria may be inhibited by either nuclease resistant or nuclease sensitive oligonucleotides. It was also well known that methylcarbamate modified oligonucleotides (the methylcarbamate replaced the
- 20 phosphodiester bonds) of three and four nucleotide units, and methylphosphonates longer than four nucleotide units could not enter Escherichia coli cells (Jayaraman et al., Proc. Natl. Acad. Sci. USA 78:1537-1541 (1981), Rahman et al., Antisense Res. Devel. 1:319-327 (1991)). Thus, the prior art
- 25 teaches that the alleged results described in the '533 patent conflict with previously reported results from bacterial experiments using nuclease resistant oligonucleotides, or phosphodiester oligonucleotides.

In 1993, Chrisey reported uptake in vitro of

30 phosphorothioate oligonucleotides into Vibrio bacteria under hypertonic conditions, and were only able to show uptake when the cells were grown under conditions that enhanced the permeability of the bacterial cells (i.e., in a hypertonic minimal medium). From these data, Chrisey et al. concluded that, in enriched media (blood, serum, and other extracellular fluids), oligonucleotides may not be preferred antibacterial agents for use in vivo.

1.5. Oligonucleotides As Antibacterial Agents

As discussed above, essentially five publications have addressed the possibility of using oligonucleotides to inhibit bacterial growth. Four out of five of these 5 publications (Rahman, Chrisey, Jayaraman, and Gasparro) teach that oligonucleotides are not able to inhibit the growth of unmodified (intact) bacteria. Additionally, the last reference (Lupski) provides no teaching of how to inhibit the growth of intact bacteria, and provides no illustrative 10 examples that such inhibition is indeed possible.

Taken as a whole, the above publications would have not provided a reasonable expectation that one could in fact use oligonucleotides to inhibit the growth of intact bacteria. The inadequacies of the background art may be explained by

- 15 the fact that the present applicants have discovered that at least several features of the design, preparation, and use of oligonucleotides may affect antibacterial efficacy. These features include, but are not limited to: 1) the dose of oligonucleotide; 2) the length of the oligonucleotide; 3) the
- 20 growth conditions used during the *in vitro* assay; 4) the chemical backbone of the oligonucleotide; and 5) the method of post-synthesis purification. Each of these features are discussed in greater detail below.

The dose of oligonucleotide may significantly effect the 25 observed amount of growth inhibition. Fig. 1 shows that the percent of inhibition varies from 100% down to about 19% as the dose of oligonucleotide is reduced from 285 μ M to 5 μ M in a standard MIC assay (described in Section 4.5, infra). Of the background references, only Rahman and Jayaraman taught 30 concentrations of oligonucleotide to the first section 4.5.

30 concentrations of oligonucleotide that fall within the disclosed range (but observed little to no inhibitory effect against intact bacteria).

The applicants have also found that the length of the oligonucleotide is directly related to its ability to

35 specifically bind and inhibit the normal function of the target sequence. Shorter oligonucleotide sequences generally have a reduced Tm (duplex melting temperature) and are thus

more likely to cause undesirable side effects of nonspecific binding or have no effect. Gao et al., Molec. Pharm. 41:223-229 (1992) have shown that, using an in vitro enzymatic assay, the inhibitory effect of an oligonucleotide sequence increased to the last of the last

- 5 increased as the length of the oligonucleotide was progressively increased from a 7mer up to a 28mer. Gao et al. observed no specific inhibitory activity when a 7mer was tested. Of the cited references, Rahman, Jayaraman, Gasparro, and Chrisey used oligonucleotides that were a
- 10 maximum of only 12 bases in length. Typically, oligonucleotides as short as the disclosed 12mers show a high degree of nonspecific binding. Lupski chose sequences of about 25 bases in length but the majority of the disclosed sequences comprised a high degree of degeneracy which allows
- 15 for binding to multiple target sites. For example, oligonucleotides comprising bases such as inosine, or "N" (which indicates the use of A, C, G, or T), are usually produced when one wishes to allow binding to sequences where the precise target sequence is unknown (Ohtsuka et al., J.
- 20 Biol. Chem. 260:2605 (1985)). Sequences with such broad based homology run the risk of nonspecific binding to host sequences and associated toxicity effects. Additionally, Lupski's teaching is inherently suspect given that no data demonstrating the inhibition of bacterial growth was
 25 provided.

It should also be noted that shorter oligonucleotide sequences generally have reduced Tm's. The oligonucleotides taught by Rahman, Jayaraman, Gasparro, and Chrisey were generally so short that the Tm's for the oligonucleotide-

- 30 target sequence hybrids were usually below 37° C. For example, the 12mer phosphorothioate sequence taught by Chrisey has a predicted Tm of 28.9° C, the 9mer taught by Gasparro had a predicted Tm of 24.7° C, and the 7mer (AGGAGGT) taught by Jayaraman and 4mer (GGAG) taught by
- 35 Rahman both had a predicted Tm's well below 10° C. Given these data, it is clear that oligonucleotides of the length

taught by these references are generally not useful as antisense or antigene agents under physiologic conditions.

The growth rate and conditions under which antibiotic susceptibility are measured may profoundly effect a

5 bacterium's sensitivity to antibacterial agents (Arrow et al., Antimicrob. Agents Chemother. 26:507 (1984)), and the uptake of the antibiotic into the cell (Arrow et al., Microbiol. Rev. 51:439-457 (1987)). Accordingly, methods for screening oligonucleotides in vitro for antibacterial

10 activity should generally be conducted under standardized conditions that reflect the in vivo circumstances of a given pathogen such as the NCCLS MIC tests (see Section 4.5, infra). None of the background references recognized that growth conditions might effect the result of antibiotic

15 susceptibility tests, and thus none of these references assayed for the inhibition of bacterial growth using the

Among other things, the antibacterial efficacy of an oligonucleotide may be directly related to the relative

20 nuclease resistance of the chemical backbone of the oligonucleotide. Gasparro and Lupski did not recognize this facet of the present invention and thus did not teach oligonucleotides that were designed to be nuclease resistant.

Consequently, the oligonucleotides used by Gasparro and 25 Lupski would have been rapidly degraded by the cell (see Section 1.6, infra), and would thus have little utility as antibacterial agents.

Additionally, the post-synthesis handling and purification of the oligonucleotides may profoundly effect 30 antibacterial efficacy. None of the background references recognized the particular importance of post-synthesis handling, and thus none of the references explicitly suggest or describe purification protocols that produce effective antibacterial oligonucleotides.

In summary, none of the background references recognized the importance of the features described above. In brief, Rahman and Jayaraman both failed to provide explicit teaching

of oligonucleotides of the correct length, the use of proper susceptibility assays, or the correct purification scheme; Gasparro failed to explicitly teach the correct dose of oligonucleotide, oligonucleotides of the correct length, the

- 5 use of proper susceptibility assays, the importance of nuclease resistant backbones, or the correct purification scheme; Chrisey failed to explicitly teach the correct dose of oligonucleotide, oligonucleotides of the correct length, the use of proper susceptibility assays, or the correct
- 10 purification scheme; and Lupski failed to explicitly teach the correct dose of oligonucleotide, the use of proper susceptibility assays, the importance of nuclease resistant backbones, or the use of purified oligonucleotides. The background references, considered as a whole, failed to
- 15 recognize the importance of all of the features described above. Furthermore, none of the background references used intact clinical isolates for their studies. Accordingly, the use of oligonucleotides to inhibit the growth of clinically relevant (i.e., intact) strains of bacteria remained elusive.
- 20 Conversely, the present disclosure teaches the importance of all of the above features, and integrates all of them to provide the <u>first</u> teaching of the use of antibacterial oligonucleotides to inhibit the growth of clinically relevant bacterial pathogens.

25

1.6. Nuclease Resistant Oligonucleotides

It has been demonstrated that the fate of internalized oligonucleotides is critical to the success of antisense gene therapy (Bennett, Antisense Res. Devel. 3:235-241 (1993)).

- 30 The rapid intracellular degradation of oligonucleotides is a barrier to efficient inhibition of gene expression. One of the major problems in utilizing naturally occurring phosphodiester oligonucleotides is their rapid degradation by nucleases in mammalian cells or in serum-containing culture
- 35 medium (Cohen, <u>Oligodeoxynucleotides</u>: <u>Antisense Inhibitors</u>
 of <u>Gene Expression</u>, Boca Raton, Fla., CRC Press (1989)).
 There is abundant evidence that modification of the backbone

of oligonucleotides confers varying degrees of nuclease resistance. Hoke et al., Nucl. Acids Res. 19:5743 (1991) compared phosphodiester backbone oligonucleotides to fully modified phosphorothicate backbone oligonucleotides, and to chimeric phosphodiester and phosphorothicate backbone oligonucleotides. Hoke et al. demonstrated that the phosphorothicate oligonucleotides were degraded up to 45 times slower than the phosphodiester or chimeric backbone oligonucleotides.

- There have been reports that chimeric oligonucleotides that are end-capped with nuclease resistant backbone linkages are resistant to degradation (Cohen, "Oligodeoxynucleotides: Antisense Inhibitors of Gene Expression," Boca Raton, Fla., CRC Press (1989)). However, Hoke et al. teach that capped
- 15 oligonucleotides are rapidly degraded by intracellular endonucleases, and thus, that capping oligonucleotides with nuclease resistant modifications may not be sufficient for sustaining pharmacological activities of oligonucleotides in cells. Finally, Hoke et al. concludes that while capping of
- 20 oligonucleotides may provide protection from exonucleases in cell culture, the action of intracellular endonucleases is sufficient to degrade these capped oligonucleotides when they enter a cell.

Hoke et al. is corroborated by Gao et al. who studied 25 the relationship between the structure of the phosphodiester/phosphorothioate chimeras and nuclease resistance. Gao et al. showed a correlation between the number of phosphorothioate linkages and nuclease resistance of the oligonucleotide.

- Devel. 3:53-66 (1993), have looked at the effects of backbone modifications on cellular uptake of oligonucleotides in eukaryotes. This is an important property as the efficacy of an antisense oligonucleotide will be influenced by cellular
- 35 uptake. Zhao et al. demonstrated that cell surface binding and uptake was greatest for phosphorothioate oligonucleotides followed by phosphodiester/phosphorothioate chimeras, and

finally by phosphodiester backbone oligonucleotides. Chrisey et al., Antisense Res. Devel. 3:367-381 (1993), looked at the uptake and stability of phosphodiester and phosphorothicate backbone oligonucleotides by bacteria under hypertonic conditions. Chrisey et al. concluded that phosphorothicate 6mers were relatively resistant to nuclease activity in Vibrio parahaemolyticus cells and were relatively non-toxic. However, Chrisey et al. did not demonstrate that the internalized 6mers had antimicrobial activity.

- Various modifications to the oligonucleotide backbone have been found to inhibit nuclease degradation. Such nuclease resistant modified nucleotides are well described in the literature and include, but are not limited to, the methylphosphonates, p-ethoxy deoxyribonucleotides, p-ethoxy
- 15 2'-O-methyl ribonucleotides, 2'-O-methyl ribonucleotides, phosphorothioates, and others. A brief description of representative nuclease resistant oligonucleotide backbones follows:

Methylphosphonate oligonucleotides, in addition to
20 exhibiting enhanced nuclease resistance, also have increased
hydrophobicity over phosphodiester oligonucleotides and
therefore have greater permeability to cell membranes as
compared to phosphodiester or other more highly charged
oligonucleotides.

p-Ethoxy deoxyribonucleotides have an ethyl group olinked to the phosphate backbone. p-Ethoxy deoxyribonucleotides are resistant to nuclease degradation. p-Ethoxy ribonucleotides have the following structure:

30

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20

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5′

Phosphorothicates are compounds in which one of the non-bridging oxygen atoms in the phosphate backbone of the nucleotide is replaced by a sulfur atom. The

25 phosphorothioates are resistant to cleavage by nucleases and, since they have the same number of charged groups as phosphodiester oligonucleotides, have good solubility in water. These compounds also exhibit more efficient hybridization with complementary DNA sequences than the 30 corresponding methylphosphonate analogues.

Methyl carbonates are compounds in which one of the nonbridging oxygen atoms in the phosphate backbone has been replaced by a methyl carbonate group.

2'-O-methyl ribonucleotides are compounds in which the 35 2' position of the ribose sugar ring has a methoxy group in

place of the normal hydroxyl group. 2'-O-methyl ribonucleotides have the following general structure:

5'

5

10 O OMe

20

3'

OMe

Secondary structure can also be used to make
25 oligonucleotides resistant to nucleases. Oligonucleotides with a hairpin loop structure extending from the 3'-terminus, stabilizing the oligonucleotide against 3'-nucleolytic degradation, have been reported by Khan and Coulson, Nucl. Acids Res. 21(12):2957-2958 (1993). The Tm of the modified oligonucleotide from its complementary mRNA target was unaffected by the presence of the loop modification.

Further, end modification of oligonucleotides can also render an oligonucleotide resistant to nucleases, such as, for example, attaching cholesterol, psoralen, rhodamine, 35 fluorescein, DNP, amine groups, biotin, inverted (3'-3' or 5'-5') linkages, and the like, to the end of the oligonucleotide in order to render it more nuclease resistant.

2.0. SUMMARY OF THE INVENTION

The present invention relates to methods for the treatment of animals, including humans, that have a bacterial disease. The preferred method of treatment comprises the 5 administration of a purified antibacterial oligonucleotide having about 8 to about 80 nucleotides to the animal in an amount sufficient to inhibit bacterial growth, alleviate a symptom of the infection, or in an amount effective for treatment.

- The purified antibacterial oligonucleotides of the present invention will preferably bear an enhanced ability to inhibit the growth of bacterial cells relative to previously disclosed oligonucleotide preparations. The present invention also represents the first disclosure of the use of
- 15 oligonucleotides to inhibit the growth of intact clinically relevant bacteria. The oligonucleotides generally inhibit bacterial growth by acting as antisense or antigene inhibitors of bacterial gene expression (when targeted to bacterial nucleic acid sequences), or by acting aptamerically
- 20 to alter the function of specific bacterial proteins or polypeptides (when associating target amino acid sequences contained in bacterial peptides, polypeptides, and proteins). Alternatively, the oligonucleotides are targeted to an antibiotic resistance gene to render the bacteria sensitive
- 25 to a conventional antibiotic. In preferred embodiments, the antibacterial oligonucleotides are substantially nuclease resistant (i.e., resistant to nuclease activity).

Additional embodiments of the present invention are antibacterial oligonucleotides that have been produced by a 30 process that enhances the oligonucleotide's antibacterial activity. In particular, the presently described antibacterial oligonucleotides will be produced, or otherwise purified, by a process comprising either individually or in combination ion exchange or reverse phase chromatography,

35 extractions, precipitations, gel filtrations, dialysis, diafiltration or functional equivalents. Column chromatography may be by traditional of methods or High-

Performance Liquid Chromatography (HPLC), fast performance liquid chromatography (FPLC), and the like. Additionally, the oligonucleotides may be purified by processes including, for example, extraction or precipitation with alcohols or 5 organic solvents.

The present invention further contemplates the use of the described antibacterial oligonucleotides, in conjunction with an acceptable pharmaceutical carrier, to prepare medicinal compositions for the treatment of bacterial infections in animals, and more preferably mammals, including humans.

3.0. BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows a dose response curve of different

15 concentrations of antibacterial oligonucleotide NBT 89 (SEQ ID NO. 61) when tested against Escherichia coli ATCC accession No. 25922.

Figure 2 provides a nonexhaustive graph of the types of bacterial genes which proved susceptible to inhibition by 20 antibacterial oligonucleotides. The ordinate shows the categories of bacterial genes defined in Table 2(A-W).

Figures 3(a-c) show the percent inhibition of the growth of the indicated target bacteria after addition of the indicated oligonucleotide as a function of time.

Figures 4(a-c) show the percent inhibition of the growth of the indicated target bacteria after addition of the indicated oligonucleotide as a function of time.

Figures 5(a and b) show the percent inhibition of the growth of the indicated target bacteria after addition of the 30 indicated oligonucleotide as a function of time.

Figures 6(a-t) are plots of log bacterial growth (and accompanying control cultures) as a function of time after the addition of the indicated oligonucleotide (i.e., "NBT 114" indicates oligonucleotide sequence 114 (SEQ ID NO. 112)

35 from Table 1, infra). A clinical isolate of Escherichia coli ATCC accession No. 35218 (multiple drug resistant) was used in the experiments corresponding to figures 6(a-t).

Figures 7(a-j) are plots of log bacterial growth (and accompanying control cultures) of the penicillin resistant clinical isolate of *Staphylococcus aureus* ATCC accession No. 13301 as a function of time after the addition of the 5 indicated oligonucleotide.

Figures 8 shows that animals challenged with the bacterial pathogen *Escherichia coli* show a significant increase in survival after treatment with oligonucleotide 114 (SEQ ID NO. 112) relative to nontreated control animals.

- Figure 9 shows that test animals infected with the bacterial pathogen Staphylococcus aureus show a significant increase in survival after treatment with the variant of oligonucleotide 114 (SEQ ID NO. 112), SOT 114.21, relative to nontreated control animals.
- Figures 10(a-b) show the results observed when the indicated antibacterial oligonucleotides were tested for bactericidal activity against *Staphylcoccus aureus* using a standard overnight MIC assay.

Figures 11(a-b) show the results observed when the 20 indicated antibacterial oligonucleotides were tested for bactericidal activity against Serratia liquefaciens using a standard overnight MIC assay.

Figure 12 shows the results obtained when the indicated antibacterial oligonucleotides were tested using a standard 25 MIC assay against Staph. aureus.

Figure 13 shows the results obtained when a variety of different length versions of the indicated antibacterial oligonucleotide were tested using a standard MIC assay against *Staph. aureus*.

Figure 14 shows the results obtained when drug sensitive and drug resistant Staph. aureus were treated with oligonucleotide 114, and ampicillin.

Figure 15 shows the results of a standard MIC assay using oligonucleotide MMT 114.15 against *P. aeroginosa* strain 35 10145.

Figure 16 shows the results of a standard MIC assay using SOT 114.21 against Strep. pyogenes strain 14289.

4.0. DETAILED DESCRIPTION OF THE INVENTION

Prior to the present invention, clinically relevant bacterial pathogens were largely immune from treatment with antisense oligonucleotides. The reasons that the prior art oligonucleotides were ineffective against these pathogens include the dosages used, the lack of nuclease resistance of the oligonucleotide or the choice of the backbone, the length of the oligonucleotide, and the method of purification.

The present invention describes a method for generating 10 oligonucleotides having the novel property of being capable of having bacteriostatic or bactericidal effects on clinically relevant bacterial pathogens. The oligonucleotides generated using the presently described methods are contemplated to be able to exert antibacterial

- 15 effect both in vitro and in vivo. Typically, the antibacterial oligonucleotides will be targeted to bacterial sequences where, after associating with or binding to the target sequence, the oligonucleotide disrupts the normal function of the target sequence. The antibacterial effect of
- 20 the oligonucleotide may be caused by either specific or nonspecific association as long as bacterial growth is inhibited.

Accordingly, particularly preferred embodiments of the present invention include the novel antibacterial

25 oligonucleotides, methods of making the antibacterial oligonucleotides, and methods of using the novel antibacterial oligonucleotides to treat bacterial infection.

Given that bacterial infection is a particularly problematic complication in immunocompromised individuals

30 such as patients suffering from acquired immunodeficiency disease syndrome (AIDS), HIV infected individuals, patients undergoing chemotherapy or radiation therapy, etc., an additional embodiment of the presently described invention is the use of the presently described antibacterial

35 oligonucleotides to treat immunocompromised patients.

In a particularly preferred embodiment, the antibacterial oligonucleotides may be used to treat bacterial

infections in conjunction with similarly engineered antiviral oligonucleotides that are directed to any of a wide variety. of human viruses including, but not limited to, adenovirus, human immunodeficiency virus, human leukemia virus, rhino 5 virus, herpes virus, human papilloma virus, respiratory syncytial virus, cytomegalo virus, Epstein bar virus, hepatitis virus (A, B, C and delta), etc. Accordingly, an additional embodiment of the presently described invention are mixed oligonucleotide compositions that comprise both 10 antiviral and antimicrobial (e.g., antifungal, antibacterial, antiparasitic, etc.) oligonucleotides. Preferably, the relative ratios of the oligonucleotides present in such compositions shall be adjusted to target bacterial, parasitic, fungal, yeast, and viral pathogens that are 15 generally associated as secondary infectious sequelae of infection by one another.

An additional embodiment of the present invention are therapeutic oligonucleotides that fuse one or more sequences with known antimicrobial, antibacterial, or antiviral 20 therapeutic activity. Such fusions are deemed to constitute novel compositions having broad spectrum activity against multiple and distinct bacterial species, as well as broad antiviral and antibacterial activities. Similarly, oligonucleotides bearing multiple active sequences, or mixed 25 compositions of antibacterial oligonucleotides, may be used to target the activity of a gene product in an pathogen by blanket targeting of the DNA (via triplex inhibition, disrupting DNA replication, etc.) and RNA (via RNase H activation or directly disrupting translation, etc.) encoding 30 the activity of interest, as well as by aptameric inhibition of the gene product.

Where the therapeutic use of the presently described antibacterial oligonucleotides is contemplated, the antibacterial oligonucleotides are preferably administered in 35 a pharmaceutically acceptable carrier, via oral, intranasal, rectal, topical, intraperitoneal, intravenous, intramuscular, subcutaneous, intracranial, subdermal, transdermal,

intrathecal methods, or the like. Typically, the preferred formulation for a given antibacterial oligonucleotide is dependent on the location of the target organism in the host animal or the location in a host where a given infectious 5 organism would be expected to initially invade.

For example, topical infections are preferably treated or prevented by formulations designed for topical application, whereas systemic infections are preferably treated or prevented by administration of compositions

10 formulated for parenteral administration. Additionally, pulmonary infections may be treated both parenterally and by direct application of the antibacterial oligonucleotides to the lung by inhalation therapy.

Additionally, as oligonucleotides are cleared from the 15 bloodstream, they can often accumulate at relatively high levels in the kidneys, liver, spleen, lymph glands, adrenal gland, aorta, pancreas, bone marrow, heart, and salivary glands. Oligonucleotides also tend to accumulate to a lesser extent in skeletal muscle, bladder, stomach, esophagus,

- 20 duodenum, fat, and trachea. Lower still concentrations are typically found in the cerebral cortex, brain stem, cerebellum, spinal cord, cartilage, skin, thyroid, and prostate (see generally Crooke, 1993, Antisense Research and Applications, CRC Press, Boca Raton, FL). Interestingly,
- 25 pathogenic bacteria also tend to accumulate in many of the above organs. Consequently, the presently described antibacterial oligonucleotides can be used to target bacterial infections in specific target organs and tissues.
- One of ordinary skill will appreciate that, from a

 30 medical practitioner's or patient's perspective, virtually
 any alleviation or prevention of an undesirable symptom
 (e.g., symptoms related to the presence of bacteria in the
 body) would be desirable. Thus, the terms "treatment",
 "therapeutic use", or "medicinal use" used herein shall refer

 35 to any and all uses of the claimed antibacterial
 - oligonucleotides which remedy a disease state or symptoms, or otherwise prevent, hinder, retard, or reverse the progression

of disease or other undesirable symptoms in any way whatsoever.

Preferably, animal hosts that may be treated using the oligonucleotides of the present invention include, but are 5 not limited to, invertebrates, vertebrates, birds (such as chickens and turkeys, etc.) fish, mammals such as pigs, goats, sheep, cows, dogs, cats, and particularly humans.

When used in the therapeutic treatment of disease, an appropriate dosage of an antibacterial oligonucleotide, or 10 mixture thereof, may be determined by any of several well established methodologies. For instance, animal studies are commonly used to determine the maximal tolerable dose, or MTD, of bioactive agent per kilogram weight. In general, at least one of the animal species tested is mammalian. Those 15 skilled in the art regularly extrapolate doses for efficacy and avoiding toxicity to other species, including human. Before human studies of efficacy are undertaken, Phase I clinical studies in normal subjects help establish safe doses. Additionally, therapeutic dosages may also be altered 20 depending upon factors such as the severity of infection, and the size or species of the host.

The presently described antibacterial oligonucleotides may also be complexed with molecules that enhance their ability to enter the target cells. Examples of such 25 molecules include, but are not limited to, carbohydrates, polyamines, amino acids, peptides, lipids, and molecules vital to bacterial growth.

Additionally, the antibacterial oligonucleotide may be complexed with a variety of well established compounds or 30 structures that, for instance, further enhance the *in vivo* stability of the oligonucleotide, or otherwise enhance its pharmacological properties (e.g., increase *in vivo* half-life, reduce toxicity, etc.).

The use of synthetic oligonucleotides are advantageous 35 as an approach to treating bacterial infection because sequences can be specifically designed to inhibit bacterial

growth while not interfering with the metabolism of mammalian cells.

The present invention also relates to oligonucleotides that have demonstrated antibacterial activity in vitro. In 5 particular, the oligonucleotides will have antibacterial activity as measured in a MIC (minimal inhibitory concentration) test that is recognized in the art as predictive of in vivo efficacy for the treatment of a bacterial infection with antibiotics. Without pretreatment 10 of the bacteria to permeabilize them and without PEG-

- 10 of the bacteria to permeabilize them and without PEGmodification of the oligonucleotides, the oligonucleotides of
 the present invention are able to hybridize to a targeted
 region of a chosen bacterial polynucleotide (DNA or RNA) to
 effectively inhibit the ability of that polynucleotide to
- 15 serve as a template for synthesis of its encoded product (DNA, RNA or protein), or otherwise inhibit the target sequence's normal function in the bacterium, thereby causing a bacteriostatic or bactericidal effect. Certain oligonucleotides may exert their bacteriostatic or
- 20 bactericidal effects through binding to and inhibition of protein (aptameric effects).

In a preferred embodiment, the invention uses oligonucleotides that are substantially nuclease resistant. This includes oligonucleotides completely derivatized by

- 25 phosphorothioate linkages, 2'-0-methylphosphodiesters, pethoxy oligonucleotides, peisopropyl oligonucleotides, phosphoramidates, chimeric linkages, and any other backbone modifications, as well as other modifications, which render the oligonucleotides substantially resistant to endogenous
- 30 nuclease activity. Additional methods of rendering an oligonucleotide nuclease resistant include, but are not limited to, covalently modifying the purine or pyrimidine bases that comprise the oligonucleotide. For example, bases may be methylated, hydroxymethylated, or otherwise
- 35 substituted (glycosylated) such that the oligonucleotides comprising the modified bases are rendered substantially nuclease resistant.

The present invention further relates to compositions comprising nuclease resistant antibacterial oligonucleotides. These compositions generally comprise the oligonucleotide (or a mixture of oligonucleotides) and a physiologically 5 acceptable carrier. After administration, the oligonucleotides enter the bacterial cell and bind to the target. The target may be a polynucleotide where hybridization to the oligonucleotide results in an inability of the polynucleotides to serve as templates for their 10 encoded products. When the target is a protein, the bound oligonucleotide protein complex is inhibited relative to normal protein function (aptameric effect). As a result, growth of the bacteria are inhibited and the effects of the bacteria on the animal are less than they would have been if 15 the oligonucleotides had not been administered.

Optionally, the presently described antibacterial oligonucleotides may be formulated with a variety of physiological carrier molecules. For example, the antibacterial oligonucleotides may be combined with a lipid (or cationic lipid), the resulting oligonucleotide/lipid emulsion, or liposomal suspension may, inter alia, effectively increase the in vivo half-life of the oligonucleotide. The use of cationic, anionic, and/or neutral lipid compositions or liposomes is generally described in International Publications Nos. WO 90/14074, WO 91/16024, WO 91/17424, Pat. No. 4,897,355, herein incorporated by reference.

The antibacterial oligonucleotides of the present invention may also be introduced into bacteria after being 30 complexed with cationic lipids such as DOTMA (which may or may not form liposomes) which complex is then contacted with the target cells. Suitable cationic lipids include, but are not limited to, N-(2,3-di(9-(Z)-octadecenyloxyl))-prop-1-yl-N,N,N-trimethylammonium (DOTMA) and its salts, 1-O-oleyl-2-O-oleyl-3-dimethylaminopropyl-β-hydroxyethylammonium and its salts and 2,2-bis (oleyloxy)-3-(trimethylammonio) propane and its salts. By assembling the antibacterial oligonucleotides

into lipid-associated structures, the antibacterial oligonucleotides may be targeted to specific bacterial cell types by the incorporation of suitable targeting agents (i.e., specific antibodies or receptors) into the 5 oligonucleotide/lipid complex.

In another embodiment, the presently described purified oligonucleotides may be complexed with additional antibacterial agents. Additionally, the described nuclease resistant antibacterial oligonucleotides may also be linked to a conventional antibiotic or other chemical group that inhibits bacterial gene expression.

Having a demonstrated activity in vitro, the presently described antibacterial oligonucleotides are also contemplated to be effective in compating bacterial

15 contamination of laboratory cultures, consumables (food or beverage preparations), or industrial processes.

4.1. <u>Definitions</u>

- For the purposes of the present disclosure, the term
 20 "oligonucleotide" typically refers to a molecule comprising
 from about 8 to about 80 nucleotides, preferably about 15 to
 about 35 nucleotides, including polymers of ribonucleotides,
 deoxyribonucleotides, or both, with the ribonucleotide and/or
 deoxyribonucleotides being connected together via 5' to 3'
- 25 linkages that may include any of the linkages known in the oligonucleotide art (including, for example, oligonucleotides comprising 5' to 2' linkages). In general, longer oligonucleotides (about 50 nucleotides) display enhanced targeting specificity but may be less efficient gaining entry
- 30 to the target bacterium. Conversely, shorter oligonucleotides may more easily permeate the target bacteria, but may display a tendency to nonspecifically associate with host sequences and create a bystander effect or have no effect at all. Additionally, shorter
- 35 oligonucleotides may less efficiently bind to, and thus nonspecifically inhibit, bacterial target sequences. For example, shorter antisense oligonucleotides (6mers to 7mers)

may prove less efficient at specifically binding the target mRNA, and may prove less efficient at activating RNase H activity. Shorter oligonucleotides may also effect host gene expression in a nonspecific, and thus undesirable, manner.

- In spite of the above, the present application additionally contemplates relatively short oligonucleotide sequences (6mers to 7mers) having the desired antibacterial effects, and preferably broad-spectrum antibacterial effects, while exhibiting few adverse side effects in the host. In
- 10 fact, an example of a short (6mer) oligonucleotide is provided below that exhibits significant antibacterial activity and is contemplated as a specific example of a preparation of an antibacterial oligonucleotide that functionally defines the lower size limit of the present
- 15 invention. Given that the present invention specifically contemplates short oligonucleotides with demonstrated antibacterial function, the short oligonucleotides of the present invention specifically exclude short inoperative oligonucleotides such as AGGAGGT or GGAG.
- Accordingly, additional embodiments of the present invention include relatively short (e.g. 6mers) oligonucleotides that have been identified by using the presently disclosed methods of synthesis in conjunction with standard antibacterial assays while gradually deleting bases from oligonucleotides with established antibacterial activity in order to define short antibacterial "core" sequences.

A particular embodiment of the present application contemplates oligonucleotides that have been modified to enhance the specificity of binding. Increased specificity 30 allows for shorter oligonucleotides having the desirable features of both long and short oligonucleotides.

The presently described oligonucleotides may be constructed using either conventional bases (adenosine, cytosine, guanosine, thymidine, xanthine, inosine, or 35 uridine) or any other modified bases, or base analogues that allow an oligonucleotide comprising such analogues to retain its ability to hybridize to a complementary nucleotide

sequence. Examples of such non-naturally occurring bases that are capable of forming base-pairing relationships with naturally occurring nucleotide bases include, but are not limited to, aza and deaza pyrimidine analogues, aza and deaza purine analogues as well as other heterocyclic base analogues, wherein one or more of the carbon and nitrogen atoms of the purine and pyrimidine rings have been substituted by heteroatoms, e.g., oxygen, sulfur, selenium, phosphorus, and the like.

- .10 Modified oligonucleotides, nuclease resistant oligonucleotides, and antisense oligonucleotides are also meant to be encompassed by this definition. The term "oligonucleotide" is meant to encompass all of the foregoing, unless the context dictates otherwise.
- The term "modified oligonucleotide" refers to oligonucleotides that include one or more modifications of the nucleic acid bases, sugar moieties, internucleoside phosphate linkages, as well as molecules having added substituents, such as diamines, cholesteryl or other
- 20 lipophilic groups, or a combination of modifications at these sites. The internucleoside phosphate linkages can be phosphodiester, phosphotriester, phosphoramidate, siloxane, carbonate, carboxymethylester, acetamidate, carbamate, thioether, bridged phosphoramidate, bridged methylene
- 25 phosphonate, bridged phosphoramidate, bridged phosphoramidate, bridged methylene phosphonate, phosphorothioate, methylphosphonate, phosphorodithioate, bridged phosphorothioate and/or sulfone internucleotide linkages, or 3'-3' or 5'-5' linkages, and combinations of
- 30 such similar linkages (to produce mixed backbone modified oligonucleotides). The modifications can be internal or at the end(s) of the oligonucleotide molecule and can include additions to the molecule of the internucleoside phosphate linkages, such as cholesteryl, diamine compounds with varying
- 35 numbers of carbon residues between amino groups and terminal ribose, deoxyribose and phosphate modifications which cleave or cross-link to the opposite chains or to associated enzymes

or other proteins. Electrophilic groups such as ribosedialdehyde could covalently link with an epsilon amino group of the lysyl-residue of such a protein. A nucleophilic group such as n-ethylmaleimide tethered to an oligomer could 5 covalently attach to the 5' end of an mRNA or to another electrophilic site. The term modified oligonucleotides also includes oligonucleotides comprising modifications to the sugar moieties such as 2'-substituted ribonucleotides, or deoxyribonucleotide monomers, any of which are connected 10 together via 5' to 3' linkages. The term "modified oligonucleotide" is meant to encompass all of the foregoing, unless the context dictates otherwise, and also refers to oligonucleotides comprising chemical groups (e.g., sugar molecules, amino acids, etc.) that may improve the 15 antibacterial activity of the oligonucleotide.

The term "oligonucleotide backbone" refers to any and all means of chemically linking nucleotides such that oligonucleotides result that are capable of base-pairing or otherwise hybridizing, or interacting with a bacterial target 20 sequence in a more-or-less sequence specific manner.

The term "purified oligonucleotide" refers to an oligonucleotide that has been isolated so as to be substantially free of, inter alia, incomplete oligonucleotide products produced during the synthesis of the desired 25 oligonucleotide. Preferably, a purified oligonucleotide will also be substantially free of contaminants which may hinder or otherwise mask the antibacterial activity of the oligonucleotide. In general, where an oligonucleotide is able to bind to, or gain entry and inhibit the growth of a 30 bacteria, it shall be deemed as substantially free of contaminants that hinder antibacterial activity. One example of a method to produce such purified oligonucleotides is described herein. In particular, an oligonucleotide preparation shall generally be considered substantially free 35 of adverse contaminants (e.g., contaminants that hinder the measured antibacterial activity of the nucleotides such as

alkyl amines, alkyl ammonium groups, or agents that block

oligonucleotide entry, etc.) when the sample proves effective in an in vitro MIC assay to an extent that is displays more than about twice, and preferably about five times, and most preferably at least about an order of magnitude greater

5 antibacterial activity than a corresponding preparation that has not been treated to remove the adverse contaminants. Typically, an oligonucleotide preparation shall preferably be considered substantially free of adverse contaminants when the levels of contaminants in a sample are reduced to about 1/20th of the levels found in unpurified (or intermediately purified) samples, more typically about 1/50th of the levels found in unpurified samples than about 1/100th of the levels found in intermediately or unpurified samples of oligonucleotide.

- Alternatively, an antibacterial oligonucleotide preparation may generally be considered free of adverse contaminants when the composition is about 95 percent free, and specifically about 99 percent free of contaminating alkyl amines, alkyl ammonium groups, or a mixture thereof as
- 20 compared to unpurified crude or intermediately purified samples of the oligonucleotide preparation (as measured by conductivity, mass spectroscopy, or the extent to which a given oligonucleotide preparation retains antibacterial activity).
- The term "substantially nuclease resistant" refers to oligonucleotides that are resistant to nuclease degradation, as compared to unmodified oligonucleotides, and include, but are not limited to oligonucleotides with modified backbones, such as, for example, phosphorothioates, methylphosphonates,
- 30 ethylphosphotriesters, 2'-O-methylphosphorothioates, 2'-O-methyl-p-ethoxy ribonucleotide, 2'-O-methyl ribonucleosides, methyl carbamates, and methyl carbonates, inverted bases or chimeric versions of these backbones. Typically, the relative nuclease resistance of an oligonucleotide will be
- 35 measured by comparing the percent digestion of a resistant oligonucleotide with the percent digestion of its unmodified counterpart (i.e., a corresponding oligonucleotide with

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"normal" backbone, bases, and phosphodiester linkage). nuclease resistance tests generally add a given concentration of oligonucleotide (e.g., about 121 μ molar) to a given amount of nuclease S1 (at about 0.05 units per ml final 5 concentration in the reaction), P1 (at about 0.05 units per ml final concentration in the reaction), SVP (at about 0.05 units per ml final concentration in the reaction), Micrococcal Nuclease (at about 0.5 units per ml final concentration in the reaction), etc., and measure the percent 10 degradation (all reactions are incubated at about 37°C in the buffer appropriate for each nuclease. For example, S1 nuclease digestion conditions are typically 30 mM sodium acetate (pH 4.5), 50 mM NaCl, 1 mM ZnCl2, 5% Glycerol; Pl nuclease digestion conditions are typically 30 mM sodium 15 acetate (pH 5.3), 0.2 mM ZnCl2; SVP digestion conditions were 100 mM Tris (pH 8.9) 100 mM NaCl, 14 mM MgCl2; and Micrococcal nuclease digestion conditions are typically 50 mM sodium borate (pH 8.8), 5 mM NaCl, 2.5 mM CaCl₂). Percent degradation may be determined by using analytical HPLC to 20 assess the loss of full length oligonucleotide, or by any other suitable methods (e.g., by visualizing the products on a sequencing gel using staining, autoradiography, fluorescence, etc., or measuring a shift in optical density).

Degradation is generally measured as a function of time.

25 Generally, a substantially nuclease resistant oligonucleotide will be at least about 25% more resistant to nuclease degradation than an unmodified oligonucleotide with a corresponding sequence, typically at least about 50% more resistant, preferably about 75% more resistant, and more 30 preferably at least about an order of magnitude more

resistant after 15 minutes of nuclease exposure.

The term "targeted to a bacterial sequence" refers to the fact that the presently described antibacterial oligonucleotides are substantially homologous, otherwise complementary, or capable of associating with a target bacterial sequence. By associating with the target bacterial sequence, the presently described antibacterial

oligonucleotides are able to disrupt or inhibit the normal function of the target sequence, and hence inhibit bacterial cell division. In general, the antibacterial oligonucleotides will associate or bind to the target 5 bacterial sequence and inhibit the function of the sequence by an antisense mechanism, an antigene (triplex) mechanism, or by stearic hindrance. Furthermore, the oligonucleotides can function through an aptameric mechanism by binding to nucleic acid binding proteins. For the purposes of the 10 present invention, the term "aptamer" shall refer to oligonucleotides that are capable of binding or otherwise interacting with peptides, polypeptides, or proteins in a manner that effects the normal function of the peptide, polypeptide, or protein.

- In order for the presently described antibacterial oligonucleotides to recycle their antibacterial activity, the oligonucleotides will generally associate with bacterial target sequences with an avidity sufficient to elicit an antibacterial effect, yet weak enough to allow the
- 20 oligonucleotide to disassociate from the reaction products (e.g., after messenger degradation, etc.) and subsequently target another molecule. One method of reducing the binding avidity, or relaxing the binding specificity, of an oligonucleotide is to truncate, or delete, a portion of the 25 oligonucleotide.

Alternatively, another method of relaxing the binding avidity of an oligonucleotide comprises engineering a percentage of miss-match (or more-or-less neutral match, e.g., G-U base pairs) into the antibacterial nucleotide 30 sequence. By reducing the net homology of a sequence, one effectively allows for antibacterial activity while increasing the kinetics of disassociation. Accordingly, an additional embodiment of the presently claimed methods and

oligonucleotides are relaxed-specificity antibacterial

35 oligonucleotides which comprise sequence miss-matches (with
the corresponding target sequence) of up to about 60 percent,
often about 35 percent, and preferably about 20 percent, or

less. In spite of the percentage miss-match, the relaxedspecificity oligonucleotides remain capable of associating
with bacterial target sequences under physiological
temperatures and conditions. For the purposes of the present
invention, the term "miss-match" shall apply to all Watson
and Crick polynucleotide base-pairs, other than A:T, G:C, and
A:U, and the inverses thereof.

Furthermore, one of ordinary skill will appreciate that the maximally tolerated percentage miss-match may vary

10 depending on the G/C content of the oligonucleotide. In general, an A/T-rich sequence may tolerate a fairly high percentage of miss-match where the G/C base pairs have been retained. In any event, the amount of sequence miss-match should not be such that undue side effects result in the 15 host.

Additionally, given the reduced charge associated with oligonucleotides comprising partially or fully substituted chemical backbones, it is to be understood that such oligonucleotides may retain the ability to bind target

20 bacterial sequence under physiological conditions although comprising a greater amount of sequence miss-match than may be tolerated by conventional oligonucleotides.

An additional embodiment of the present invention is antibacterial oligonucleotides that are capable of inhibiting 25 bacterial growth by cross reacting with a variety of both known and unknown bacterial target sequences. For the purposes of the present disclosure, the term "cross reactive antibacterial oligonucleotide" shall refer to an oligonucleotide sequence that inhibits bacterial growth by 30 interacting with bacterial sequences that may share less than 100 percent sequence homology, and preferably at least about 50 percent sequence homology, with the oligonucleotide. Examples of such a cross reactive antibacterial activity include: instances where heterologous, similar, and 35 homologous bacterial sequences are bound and affected by an oligonucleotide that is targeted to a related sequence; instances where an antibacterial oligonucleotide is able to

interact with bacterial sequences that share a sufficient percentage of otherwise random sequence complementarity (e.g., short, interspersed regions of high sequence complementarity, etc.) with the oligonucleotide such that

- 5 bacterial growth is inhibited; and instances where a given antibacterial oligonucleotide is able to inhibit bacterial growth although all or some of the affected bacterial target sequences are unknown (this includes instances where the cross reactive oligonucleotide has up to 100% homology with
- 10 an unknown target DNA sequence). Target sequences comprised within conserved or related control regions, which are often noncoding, are deemed to constitute particularly effective targets for cross reactive antibacterial oligonucleotides that operate via an antigene mechanism.
- A "functional equivalent" of the sequences disclosed in the Sequence Listing shall include any oligonucleotides comprising sequence that is at least about 25 percent sequence homologous, preferably about 33 percent sequence homologous, and more preferably at least about 50 percent
- 20 homologous to any one of SEQ ID NOS. 1-176, and demonstrates at least about 30 percent, and preferably at least about 50 percent of the antibacterial activity of the corresponding oligonucleotide in the Sequence Listing when measured in an MIC assay.
- The term "bacterial sequence" includes any and all forms of DNA, RNA or amino acid polymers (or oligomers) that are present in the cell.

The term "competent cells" refers to bacterial cells that have been manipulated in culture or otherwise

30 chemically, osmotically, or thermally modified such that the cells bear an enhanced ability to internalize exogenous nucleic acid.

The term "pathogenic bacteria" refers to any and all bacteria that are, or have been, associated with clinical

35 symptoms of disease in animals, including humans. The term "wild-type" bacteria refers to a bacteria that has not been modified either chemically or genetically in any way

whatsoever (other than growth in culture medium). In particular, a "wild-type" bacteria shall not be genetically modified such that the bacteria has an enhanced permeability to macromolecules or biological polymers or oligomers.

- The term "antisense oligonucleotide" refers to an oligonucleotide that has a sequence that is substantially complementary to a target DNA or mRNA, so that the antisense oligonucleotide will hybridize in a complementary fashion to the DNA or mRNA to form a complex by Watson-Crick base
- 10 pairing. Generally, the antisense oligonucleotide will bind the complementary target sequence with an avidity, in vivo, sufficient to inhibit the normal function of target sequence.

The term "bacteriostatic oligonucleotide" refers to oligonucleotides that inhibit or retard the growth of 15 bacteria either in vitro or in vivo.

The term "bactericidal oligonucleotide" refers to oligonucleotides that directly, or indirectly, cause the death of bacteria either in vitro or in vivo.

The term "Gram negative bacteria" refers to the
20 inability of bacteria to resist decolorization with alcohol
after being treated with Gram's crystal violet stain.
However, following decolorization, these bacteria can be
readily counter-stained with safranin, imparting a pink or
red color to the bacterium when viewed by light microscopy.

- 25 This reaction is usually an indication that the bacterium's outer structure consists of a cytoplasmic membrane (inner), which is surrounded by a relatively thin peptidoglycan layer, which in turn, is surrounded by an outer membrane. Typical examples of Gram negative bacteria include, but are not
- 30 limited to, Escherichia, Salmonella, Edwardsiella, Arizona, Citrobacter, Enterobacter, Proteus, Yersinia, Klyvera, Klebsiella, Neiserria, Vibrio, Pasturella, Haemophilus, Pseudomonas, Moraxella, Eikenella, Fusobacterium, Acidominococcus, Actinobacillus, Cardiobacterium, Serratia,
- 35 Providencia, Erwinia, Tatumella, Shigella, Branhamella, Aeromonas, Francisella, Gardnerella, Alcalígenes, Kingella, Agrobacterium, Leptotrichia, Megasphaera, Capnocytophaga,

Cromobacterium, Hafnia, Morganella, Pectobacterium, Cadecea, Helicobacter, Morococcus, Pleisiomonas, Bordetella, Brucella, Achromobacter, Flavobacterium, Bacteroides, Veillonella, Streptobacillus, Pneumococcus, and Calymmatobacterium.

- The term "Gram positive bacteria" refers to the ability of bacteria to resist decolorization with alcohol after treatment with Gram's crystal violet stain, imparting a violet color to the bacterium when viewed by light microscopy. This reaction is usually an indication that the
- 10 bacterium's outer structure consists of a cytoplasmic membrane surrounded by a thick, rigid bacterial cell wall mainly comprised of peptidoglycan (murein). Typical examples of Gram positive bacteria include, but are not limited to, Aerococcus, Listeria, Streptomyces, Actinomadura,
- 15 Lactobacillus, Eubacterium, Arachnia, Mycobacterium,
 Peptostreptococcus, Staphylococcus, Corynebacterium,
 Erysipelothrix, Dermatophilus, Rhodococcus, Bifodobacterium,
 Lactobacillus, Streptococcus, Bacillus, Peptococcus,
 Micrococcus, Kurthia, Nocardia, Nocardiopsis, Rothia,
- 20 Propionibacterium, Actinomyces, Enterococcus, and Clostridia.

 Additionally, the presently described antibacterial oligonucleotides may be effective against bacteria including, but not limited to, Campylobacter, Spirillium, Borrelia, Treponema, Leptospira, Legionella, and Chlamydia.
- The term "mycobacterium" refers to any and all strains of bacteria drawn from the group comprising: Mycobacterium tuberculosis, Mycobacterium bovis, Mycobacterium leprae, Mycobacterium avium-intracellulare, Mycobacterium kansasii, Mycobacterium scrofulsceum, Mycobacterium marinum,
- 30 Mycobacterium fortuitum, Mycobacterium ulcerans,
 Mycobacterium chelonae, Mycobacterium paratuberculosis,
 Mycobacterium xenopi, Mycobacterium simiae, or other
 mycobacteria falling within the Runyon groups I-IV as
 described in Runyon, Med. Clin. North Amer. 43:273-290
- 35 (1959), or Mandell et al., 1990, <u>Principles and Practice of Infectious Disease</u> 3rd. ed., Churchill Livingstone Inc., New York, N.Y. 10036, herein incorporated by reference.

The term "MIC test" refers to a National Committee on Clinical Laboratory Standards ("NCCLS") approved test for determining the minimum inhibitory concentrations ("MIC") of bacteria by broth dilution. This term includes the use of 5 this test for determining the percent inhibition of bacterial growth by the oligonucleotides of the invention.

The term "transport" refers to the movement of the oligonucleotides of the invention from outside the bacterial cell across the bacterial cell's outer-structure and into the bacterial cell's cytoplasm.

The term "virulence factor" refers to bacterial products which contribute to the pathogenicity of a bacteria, such as, for example, antibiotic resistance factors, toxins (exo- and endo-), adherence factors that recognize host tissues,

15 extracellular receptors, bacterial iron-binding proteins, and surface modifications that allow the bacteria to escape the immune system (e.g., polysaccharide coats or capsules).

The term "labeled oligonucleotides" refers to oligonucleotides that have been modified to allow a 20 determination of the presence or amount of the oligonucleotide. Typical labels include, for example, radioisotopes, biotin, and enzymes (such as luciferase, or β -galactosidase).

The term "stringent conditions" generally refers to

25 hybridization conditions that (1) employ low ionic strength
and high temperature for washing, for example, 0.015 M

NaCl/0.0015 M sodium citrate/0.1% SDS at 50°C.; (2) employ
during hybridization a denaturing agent such as formamide,
for example, 50% (vol/vol) formamide with 0.1% bovine serum

30 albumin/0.1% Ficoll/0.1% polyvinylpyrrolidone/50 mM sodium
phosphate buffer at pH 6.5 with 750 mM NaCl, 75 mM sodium
citrate at 42°C; or (3) employ 50% formamide, 5 x SSC (0.75 M

NaCl, 0.075 M Sodium pyrophosphate, 5 x Denhardt's solution,
sonicated salmon sperm DNA (50 g/ml), 0.1% SDS, and 10%

35 dextran sulfate at 42°C, with washes at 42°C in 0.2 x SSC and
0.1% SDS. The above examples of hybridization conditions are
merely provided for purposes of exemplification and not

limitation. One of ordinary skill will appreciate that stringency may generally be reduced by increasing the salt content present during hybridization and washing, reducing the temperature, or a combination thereof. A more thorough treatise of such routine molecular biology techniques may be found in Sambrook et al., Molecular Cloning, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York, Vols. 1-3 (1989), and periodic updates thereof, herein incorporated by reference.

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4.2. Synthesis Of Oligonucleotides

The described oligonucleotides may be partially or fully substituted with any of a broad variety of chemical groups or linkages including, but not limited to: phosphoramidates,

- 15 phosphorothioates; p-ethoxy; alkyl phosphonate; 2'-O-methyl; 2' modified RNA; morpholino groups; phosphate esters; dithioates; 5' thio groups; propyne groups; or chimerics of any combination of the above groups or linkages (or analogues thereof), or any other chemical modifications that leave the
- 20 oligonucleotide capable of specifically binding to nucleic acid or protein.

Oligonucleotides, methylphosphonates, and phosphorothioates may be synthesized, using standard reagents and protocols, on an automated synthesizer utilizing methods

- 25 that are well known in the art, such as, for example, those disclosed in Stec et al., J. Am. Chem. Soc. <u>106</u>:6077-6089 (1984), Stec et al., J. Org. Chem. <u>50</u>(20):3908-3913 (1985), Stec et al., J. Chromatog. <u>326</u>:263-280 (1985), LaPlanche et al., Nuc. Acid. Res. <u>14</u>(22):9081-9093 (1986), and Fasman,
- 30 G.D. <u>Practical Handbook of Biochemistry and Molecular Biology</u>, 1989, CRC Press, Boca Raton, Florida, herein incorporated by reference.

The principal criteria for designing nuclease resistant oligonucleotides are: (1) retention of sequence-specific

35 base-pairing and triplex-forming interactions (i.e., the ability to associate with bacterial target sequence such that bacterial growth is inhibited); (2) increasing nuclease

stability; (3) ease of synthesis and purification. The most common strategies to date have involved neutralizing the charge on the phosphodiester backbone by substitution at, or replacement of, the phosphodiester moiety, conjugating

5 moieties at the 3' and/or 5' terminus, and substitutions at the 2'-position of ribose and deoxyribose. In particular, the addition of a 3'-3' or 5'-5' internucleotidic linkages at either end of the oligonucleotide, may inhibit degradation by the respective exonuclease (Seliger et al., 1991, Nucleosides and Nucleotides, 10:463-477). Additionally, several new strategies have recently emerged that utilize peptide interlinkages.

The synthesis of phosphoramidates is disclosed in Agrawal et al., Proc. Natl. Acad. Sci. USA 85:7079-7083

15 (1988). The preparation of phosphoramidates modified with several methoxyethyl phosphoramidate internucleoside linkages is disclosed in Dagle et al., Nucl. Acids Res. 18(6):4751-4757 (1990). These modified oligonucleotides are highly resistant to nucleolytic degradation and can also serve as a 20 substrate for RNase H (which degrades the RNA component of a DNA/RNA hybrid).

An approach for synthesizing formacetal linked dinucleosides is disclosed by Quaedflieg et al., Tetrahedron Lett. 33(21):3081-3084 (1992).

The synthesis and binding properties of pyrimidine oligonucleotides containing alternating modified and natural internucleoside linkages, formacetal and thioformacetal, is disclosed by Jones et al., J. Org. Chem. <u>58</u>:2983-2991 (1993). The thioformacetal modified oligodeoxynucleotides (ODN)

30 displayed high affinity and specificity for both singlestranded RNA and double-stranded DNA targets, indicating that this linkage is promising for both antisense and triplex (antigene) therapeutic applications.

The synthesis of hexanucleotide analogues containing 35 internucleotide diisopropylsilyl linkages is disclosed by Cormier and Ogilvie, Nucl. Acids Res. <u>16</u>(10):4583-4594 (1988). These oligonucleotides were not readily soluble in

water. It has been suggested that inserting terminal or internal phosphodiester groups, or highly hydrophilic groups would increase water solubility of these compounds.

The synthesis of acetamidate linked oligomers of mean 5 chain length 10-13 is disclosed by Gait et al., J. Chem. Soc., Perkin Trans. 1:1684 (1974).

The synthesis of dinucleotides and trinucleotides modified with carbamate (-OCO-NH-) bonds is disclosed by Mungall and Kaiser, J. Org. Chem. 42(4):703-706 (1977). The 10 carbamate linkage was found to be stable toward acid and base hydrolysis, as well as toward nucleases.

The synthesis of oligonucleotides with dimethylenesulfide (-CH₂-S-CH₂), -sulfoxide (-CH₂-SO-CH₂), and -sulfone (-CH₂-SO₂-CH₂) groups replacing phosphodiester linkages is reported by Schneider and Brenner, Tetrahedron Lett. 31(3):335-338 (1990); Huang et al., J. Org. Chem. 56:3869-3882 (1991); Musicki et al., Tetrahedron Lett. 32(10):1267-1270 (1991); Huang et al., Tetrahedron Lett. 33(19):2657-2660 (1992); and Reynolds et al., J. Org. Chem. 57:2983-2985 (1992).

The synthesis of 2'-O-alkyloligoribonucleotides, where the alkyl groups are methyl, butyl, allyl or 3,3-dimethylallyl is reviewed by Lamond, Biochem. Soc. Trans.

21:1-8 (1993). Oligomers comprised of the modified linkages formed stable duplexes that exhibited a higher Tm (upon binding complementary RNA) than unmodified RNA-RNA duplexes. Oligonucleotides containing the modified linkages are nuclease resistant. It was found that binding of allyl-modified oligomers to A/U rich mRNA sequences (typical of snRNAs) could be improved by incorporating the modified base 2-aminoadenine in the modified probe.

The synthesis of 2'-deoxyuridine analogues carrying an amino linker at the 1'-position of deoxyribose is disclosed by Ono et al., Bioconjugate Chem. 4:499-508 (1993). The 35 uridine analogues were incorporated into oligonucleotides and intercalating groups such as anthraquinone and pyrene derivatives that were attached to the amino group of the

linker. Several oligonucleotides were synthesized that incorporated the analogues at several different sequence positions. Duplexes formed with the analogues were more stable than unmodified duplexes. Also, the oligonucleotide analogues were resistant to exo- and endonuclease degradation. Moreover, duplexes formed with the analogues were capable of activating RNase H. The authors suggested that the bulky group attached at the C1'-position stearically masked the phosphodiester linkage from nuclease attack.

The synthesis of uniformly modified 2'-deoxy-2'-fluoro phosphorothicate oligonucleotides is disclosed by Kawasai et al., J. Med. Chem. 36:831-841 (1993). Since 2'-deoxy-2-fluororibose adopts the 3'-endo conformation, it was hypothesized that deoxy oligomers modified at the 2'-position with fluorine would adopt more uniform and more stable duplexes with RNA. The modified oligomers were found to possess thermal stabilities similar to or higher than those of the corresponding RNA duplexes. The modified oligomers demonstrated resistance to nucleases, but did not activate RNAse H.

A description of the synthesis of p-ethoxy-linked oligonucleotides may be found, *inter alia*, in application Ser. No. 08/065,016, filed May 24, 1993, herein incorporated by reference. The synthesis of inverted bases is described in Seliger et al..

Additional antibacterial oligomers may be adapted from the polynucleotide binding polymers and backbones described in Pat. Nos. 5,034,506, 5,142,047, 5,166,315, 5,185,444, 5,470,974, and 5,235,033, which are herein incorporated by reference.

The synthesis of oligonucleotides containing any of the above internucleotide linkages is well known to those skilled in the art, as is further illustrated in articles by Uhlmann et al., Chem. Rev. 90:543-584 (1990), and Schneider et al., 35 Tetrahedron Lett. 31:335 (1990). See also Reissue Pat. No. 34,069, herein incorporated by reference.

4.2.1 Oligonucleotides Comprising Modified Nucleosides

 α -Anomeric Nucleoside Units. The synthesis of a octathymidylate comprised of α -anomers is disclosed by Thuong et al., Proc. Natl. Acad. Sci. USA <u>84</u>:5129-5133 (1987). The 5 modified oligomer binds to complementary sequences containing naturally occurring β anomers. A 3'-acridine linked α -anomer was also prepared. This analogue also demonstrated sequence-specific binding. The α -anomers demonstrated nuclease stability, independently of whether linked to acridine or 10 not.

Base-Modified Nucleoside Units. The synthesis of a base analogue designed to recognize T-A and G-C Watson-Crick base pairs to facilitate sequence-specific triplex formation is disclosed by Griffin et al., J. Am. Chem. Soc. <u>114</u>:7976-7982 15 (1992).

4.3. Purification Of Oligonucleotides

The present disclosure teaches that the relative purity of an antibacterial oligonucleotide may profoundly impact its 20 antibacterial activity. As discussed in greater detail below, the antibacterial activity of an oligonucleotide may be enhanced by at least 60 percent after it has been subject to an appropriate purification protocol. It is particularly important that purification remove contaminants that either 25 obstruct the uptake of the oligonucleotides or mask the antibacterial activity of the oligonucleotides by, for example, stimulating bacterial growth.

A variety of standard methods were used to purify/produce the presently described antibacterial oligonucleotides. In brief, the antibacterial oligonucleotides of the present invention were purified by chromatography on commercially available reverse phase (for example, see the RAININ Instrument Co., Inc. instruction manual for the DYNAMAX®-300A, PureDNA™ reverse-phase columns, 1989, or current updates thereof, herein incorporated by reference) or ion exchange media (see generally, Warren and Vella, 1994, "Analysis and Purification of Synthetic

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Oligonucleotides by High-Performance Liquid Chromatography",

In Methods in Molecular Biology, vol. 26: Protocols for

Oligonucleotide Conjugates, S. Agrawal ed., Humana Press,

Inc., Totowa, NJ; Aharon et al., 1993, J. Chrom. 698:293-301;

5 and Millipore Technical Bulletin, 1992, "Antisense DNA:

Synthesis, Purification, and Analysis"). Peak fractions were

combined and the samples were desalted and concentrated by

alcohol (ethanol, butanol, isopropanol, and isomers and

mixtures thereof, etc.) precipitation, diafiltration, or gel

10 filtration followed by lyophilization, or solvent evaporation

under vacuum in commercially available instrumentation such

as, for example, a Savant Speed Vac.

Oligonucleotides of the invention were dissolved in pyrogen free, sterile, physiological saline (i.e., 0.85% 15 saline) and sterile filtered through 0.2 micron pyrogen free filters.

4.4. Oligonucleotides As Antibiotics

The principal criteria for designing antisense

20 oligonucleotides for treating bacterial infections are: (1)
retention of sequence-specific base-pairing and triplexforming interactions; (2) increasing nuclease stability; (3)
increasing the extent or kinetics of entry into the target
cell; (4) activating RNase H (while a consideration, a given
25 oligonucleotide's ability to activate RNase H is not strictly
required to observe antibacterial activity); and (5) ease of
synthesis and purification.

Although exquisite sequence specificity may be preferred in some instances, the presently described oligonucleotides

30 are capable of specifically inhibiting bacterial growth as long as they remain capable of associating with the target sequence under the relevant conditions. For example, the use of oligonucleotides to degrade RNA simply requires that the oligonucleotide associate (with at least a four base match)

35 with the bacterial RNA long enough to activate RNase H. Thus, oligonucleotides that harbor relaxed sequence specificity are deemed sufficient to activate RNase H. In

fact, because not all bacterial target sequences are known, applications are contemplated where the antibacterial oligonucleotide provides the desired inhibitory effect although not specifically targeted, or homologous, to a given 5 bacterial gene.

Modified oligonucleotides that activate RNAse H are advantageous because such oligonucleotides will hybridize to their target mRNAs and create a substrate that can be digested by RNase H. RNase H digestion destroys the target 10 mRNA, and thus, these oligonucleotides prevent the translation of the target mRNA. Accordingly, protein expression is inhibited either by the enzymatic destruction of the target mRNA, or by the oligonucleotide physically blocking translation (i.e., after the oligonucleotide 15 directly associates with ribosomal sequence).

Although RNase H activation is a factor in the design of antibacterial oligonucleotides, many antibacterial oligonucleotides (e.g., ribonucleotides targeting bacterial RNA) are not designed to activate RNase H. Typically,

- 20 modified oligonucleotides that are connected by stretches of unmodified phosphodiester linkages comprising at least about four nucleotides to about seven nucleotides should retain the ability to activate RNase H. Also, it has been observed that phosphorothicate ribonucleotides can also activate RNAse H
- 25 digestion. The differential specificity of mammalian RNase H (minimum of 5 bases) and bacterial RNAase (4 bases) affords a means of selectively targeting bacterial genes that may have strong sequence homology with certain animal genes.

Also contemplated are modified oligonucleotides that can 30 form triplexes with duplex DNA (antigene oligonucleotides), and oligonucleotides that can be used as ribozymes.

Another embodiment of the presently described antibacterial oligonucleotides is aptameric oligomers that are capable of effectively mimicking protein domains and 35 exerting an antibacterial effect by directly associating with bacterial proteins or structures.

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Additionally, antibacterial oligonucleotides may exert a therapeutic effect by specifically binding and deactivating cellular machinery. For example, the presently described oligonucleotides may directly bind ribosomal sequences and 5 inhibit translation by stearically hindering translation initiation, elongation, disassociation, or by directly destabilizing the structure of the bacterial ribosomes.

Antibiotic resistance is often caused by the presence of resistance factors that render an antibiotic ineffective. By 10 targeting resistance factors, the presently described oligonucleotides may render an otherwise antibiotic resistant organism sensitive to conventional antibiotics. Accordingly, another embodiment of the present invention is the use of antibacterial oligonucleotides in conjunction with 15 conventional antibiotics.

Another embodiment of the present invention involves the use of the presently described oligonucleotides to inhibit the expression of genes whose products regulate the replication or transfer of bacterial genes. Additionally, 20 given that antibiotic resistance genes or other virulence factors are often encoded by plasmids, antibacterial oligonucleotides targeted against plasmid replication, transfer (by conjugative transfer), or gene expression are particularly of interest. Similarly, antibacterial 25 oligonucleotides are contemplated that are capable of inhibiting the expression and transfer of genes encoded by transposable genetic elements (e.g., transposons).

4.4.1. Selection Of Targets For Oligonucleotides: <u>Gene/Operon Target Identification</u>

30

Antisense oligonucleotides which target essential structural genes, metabolic pathway genes, or transport system genes will inhibit the growth of bacterial cells. For pathogenic bacteria, virulence factors such as, for example, genes encoding antibiotic resistance, toxins, adherence and invasion factors, pili or fimbriae, flagella, antigenic variation factors, and iron binding factors, are also

preferred targets. These targets should be pathogen specific, and thus oligonucleotides directed against these targets will preferably not harm either host cells, or the normal bacterial flora of the gut.

- While some bacterial genes are expressed as individual transcripts, many are transcribed as part of a multicistronic unit or operon. Examples include the ribosomal protein operons, such as the str operon and the alpha operon in Escherichia coli. Where possible operon transcripts are
- 10 targeted. Disruption of expression of a gene in the operon may also adversely effect the expression of other genes encoded within the same operon (often in operon transcripts the translation of the 5'-most genes are required for efficient translation of the downstream genes). In theory
- 15 this could result in pleiotropic growth effects from a single oligonucleotide sequence. Specific genes and transcripts (whether expressed as part of an operon or independently) are targeted on the basis of their function in the cell. For example, the gene for glucose-6-phosphate dehydrogenase is
- 20 central to sugar metabolism. Other genes may not be relevant in our normal assay system; disruption of lactose metabolism is expected to have only a minor effect, if any, on Escherichia coli growth in media containing a more readily available carbon source such as glucose.
- Once a target gene or operon has been selected, a target region within the gene or operon sequence must be selected, for example, the start codon. An analysis of the sequences around the target sequence (e.g., 5' untranslated region, start codon, internal sequence feature, termination codon, 3'
- 30 untranslated region) is performed. This analysis generally encompasses a total of about 120 bases that flank the target sequence. This analysis further predicts the secondary structure of the antisense oligonucleotide, and can be performed using commercially available computer software.
- 35 The extended target sequence is checked for regions of stable secondary structure. The positions of the bases predicted to be involved in the stem-and-loop structures should be marked

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and the predicted Tm of the structures noted. Preferably, stem sequences should be avoided where possible. Moreover, predicted secondary structures with predicted melting temperature of 45°C or less are disregarded in this analysis.

A maximum oligonucleotide length is also selected, and the program identifies the clear regions (no stems, or the structures with the lowest melting temperatures), and also checks the loop melting temperatures for the generated oligonucleotides. Such programs are well known in the art 10 and include, for example, the program OligoTech version 1.0 (Copyright® 1995, Oligos Etc. Inc. & Oligo Therapeutics Inc.).

The length of the flanking sequence to be analyzed may be increased if an oligonucleotide with a length of greater 15 than 30 bases is selected. The transcription start site and termination site (or any attenuation sequence) are generally the most distal sequences that will be analyzed. On occasion, this may result in an analysis of about 190 or more bases of flanking sequence.

Potential oligonucleotide sequences that have high loop melting temperatures may be eliminated by the above analysis. Note that the melting temperatures for the loops obtained for the commercial programs may need to be adjusted for modified oligonucleotides since these oligonucleotides may have 25 altered base pairing avidities.

Several additional characteristics of the oligonucleotides are also considered. Stable secondary structure (potentially stable under physiologic conditions), runs of a single base (e.g., 4 or more A's), and sequences that potentially form stable homodimers are also eliminated if possible. (In cases where double-strand oligonucleotide is the desired end result, homodimers may be preferred.) The base composition of the oligonucleotide is also checked.

The two or three oligonucleotide sequences that most 35 nearly meet the above criteria are selected. Using these final oligonucleotides, the program analyzes each sequence and notes loop melting temperatures for both the sense and

the antisense strands of the candidate sequences. This decreases the possibility of the computer analysis missing a potential problem structure.

The candidate sequences, selected as above, are searched 5 for sequence matches in available sequence databases (for example, Genbank) using commercially available search software. The first search is against the bacterial sequence database(s). This allows the identification of other targets that may also be affected by the candidate sequence, and may 10 also indicate which sequences are potentially effective across bacterial genera. Since many different bacterial genera have highly related genetic organizations or related gene sequences, a potential oligonucleotide may be effective against multiple bacterial genera. For example, the 15 sequences of the gyrA genes of Escherichia coli and Salmonella typhimurium are essentially identical near the start codon.

Additionally, since bacterial translation occurs simultaneously with transcription, it may be generally 20 preferable to target antisense oligonucleotides to bacterial sequences at or near the Shine-Delgarno site (ribosome binding site) or to the translation start site of the targeted transcript.

The second search is versus a database including
25 human/primate sequences. Since these databases are still
quite limited (relative to the entire amount of sequence data
in the genome), databases generally including mammalian
sequences should be searched. Oligonucleotides that have
high specificity matches to relevant mammalian sequences
30 should be eliminated from initial consideration. (Note: that
they may be re-included after further evaluation of the
possible target sequences.)

As a consequence of the incomplete nature of the data bases comprising bacterial, primate, rodent, and mammalian 35 sequences, this method cannot ensure that all potential targets or conflicts are identified. However, as sequence data accumulates, this method will allow an experienced

practitioner of the art to identify targets and select oligonucleotide sequences for use in the methods of the invention.

5 4.5. Bacterial Inhibition Assay: MIC Test

Despite some limitations of *in vitro* susceptibility tests, the clinical data indicate that there is good correlation between MIC test results and *in vivo* efficacy of antibiotics. Murray, P., <u>Antimicrobial Susceptibility</u>

10 Testing, (Poupard et al., eds.), Plenum Press, NY, 1994;
Knudsen et al., Antimicrob. Agents Chemother. 39(6):1253-1258
(1995).

Accordingly, the presently described antibacterial oligonucleotides were tested for antibacterial activity in 15 vitro. Prior to use in vivo, a given antibacterial oligonucleotide will have demonstrated antibacterial activity in vitro against a pathogenic bacteria. Generally, the in vitro antibacterial activity of an oligonucleotide will be tested using a standard bacterial inhibition assay, or MIC test (see National Committee on Clinical Laboratory Standards "Performance Standards for Antimicrobial Susceptibility Testing" NCCLS Document M100-S5 Vol. 14, No. 16, December 1994, herein incorporated by reference).

25 4.5.1. <u>Variations On The Standard MIC Test</u>

Cells that are growing exponentially in vitro are generally not representative of cells in clinical infections where nutrients may be limited and the cells are dividing slowly or not at all, i.e., the cells are in stationary

30 phase. Starved stationary phase cells undergo a series of morphological and physiological changes that distinguish them from cells in exponential growth. These changes ensure the prolonged survival of the cells by reducing endogenous metabolism and preparing the cells for possibly adverse 35 conditions.

Further, there is a specific interrelation between the growth rate of bacterial cells and the sensitivity of the

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cells to chemicals, antibiotics, and host defenses. Thus, antibiotics developed and tested against laboratory cultures are often ineffective when directed against relatively slowly growing, clinical infections.

In an effort to address the issue of bacteria growing under starved conditions in a clinical setting, both fresh cultures and starved cultures of bacteria were used as inocula in standard MIC tests. Oligonucleotides with antibacterial activity proved effective regardless of the type of inoculum used in the MIC test.

The MIC is the lowest concentration of antimicrobial agent that completely inhibits growth of the organism in the tubes or microdilution wells as detected by the unaided eye. Viewing devices intended to facilitate reading microdilution

- 15 tests and recording of results may be used as long as there is no compromise in the ability to discern growth in the wells. The amount of growth in the wells or tubes containing the antibiotic should be compared with the amount of growth in the growth-control wells or tubes (no antibiotic) used in 20 each set of tests when determining the growth and points.
 - The percent inhibition of an oligonucleotide as reported herein was the absorbance at 625 nanometers of a bacterial culture that was treated with the oligonucleotide divided by the absorbance at 625 nanometers (i.e., O.D. 625) of a
- 25 duplicate cell culture minus oligonucleotide (control); the resulting number was subtracted from 1, and multiplied by 100%. Small variations in the optical density readings at the lower detection limit of the assay may result in calculated inhibitions of greater than 100 percent. It is
- 30 assumed that these calculations essentially represent 100 percent inhibition.

The concentration of target bacteria used in an MIC assay typically far exceeds the systemic concentrations of pathogenic bacteria that, with the possible exception of abscesses, are expected to be found in vivo. While even the presence of a single bacterium in bodily fluids is considered an indication of infection (John J. Sherris, Editor, Medical

Microbiology, An Introduction to Infectious Diseases, 2nd Edition, Elsevier, New York 1990), the precise number of bacteria/ml is not well quantified in human clinical infections (Kjeldfberg and Knight (3rd Edition), Body Fluids, 5 ASCP Press, 1993). It is difficult to quantitate bacteria in body fluids as bacteria are constantly cleared by the immune system (Myrvik, Fundamentals of Medical Bacteriology, 1974, Lea & Febiger, Publishers). In addition, bacteria grow more slowly in vivo than in vitro, so this slow growth combined 10 with the clearance by the immune system makes quantifying the number of bacteria difficult. In order to quantitate clearance of Pneumococci in the blood, Wilson (G.S. Wilson and A.A. Miles, Editors, Topley and Wilson's Principles of Bacteriology and Immunology, Williams & Wilkins, Publishers, 15 1964) reported a study where bacteria were intravenously injected into rabbits. It is evident from these data that if the immune system is unable to clear the bacteria from the blood, once the concentration of bacteria reaches 1.5 \times 10 6 cfu per ml the animal will die. In light of the above 20 discussion, the oligonucleotides need only arrest the growth of the bacteria until the immune system is capable of Furthermore, in an actual clinical situation, the clearance. concentration of bacteria/ml would be far lower than 1.5 \times 106/ml, which represents a fatal concentration in Wilson's 25 animal model.

In the presently described studies, the bacteria were grown over the period of the assays to an O.D. 600 of 0.1 as defined by the NCCLS. This represents approximately 1 x 10° concentration of bacteria which represents more bacteria/ml 30 than would be required to cause death in a clinical setting.

4.5.2. <u>Fastidious Organisms</u>

The standard media used in the MIC tests described above for the rapidly growing aerobic pathogens (Mueller-Hinton 35 medium) is not adequate for susceptibility testing of fastidious organisms. Where MIC tests are to be done using fastidious organisms, the medium, quality control procedures,

and interpretive criteria must be modified to fit each organism. For example, dilution tests for Haemophilus influenzae (using Haemophilus test medium), Nisseria gonorrhoeae (using GC agar base medium), and Streptococcus 5 pneumoniae (using lysed horse blood-supplemented, cationadjusted Mueller-Hinton broth) have been shown to be reliable It is important to note that the direct inoculum methods. suspension method of preparing the test inoculum must be used with these three species. The media and important technical 10 aspects of testing several fastidious species are described in relevant sections above and outlined in NCCLS Doc. M7-A3, Vol. 13, No. 25, entitled "Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria that Grow Aerobically -Third Edition: Approved Standard". Interpretive criteria for 15 testing these three fastidious species can also be found in NCCLS Doc. M7-A3, Vol. 13, No. 25.

4.6. Antibacterial Activity In vivo

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After demonstrating antibacterial activity in vitro, the 20 antibacterial oligonucleotide will be tested for activity in vivo. In brief, an antibacterial oligonucleotide sequence (e.g., a phosphorothioate ODN) will be tested for antibiotic activity in a mammalian test subject, and preferably a murine test subject. Phosphorothioate ODNs have previously been

- 25 tested in mammals (mice, rats, rhesus monkeys), and, when properly administered, have not been found to be significantly toxic. Prior to introduction in vivo, ODNs will be solubilized in sterile saline and serially-diluted to the desired test concentrations in sterile saline.
- Bacteria. Bacterial pathogens to be used in vivo include, but are not limited to, inter alia, the drugresistant Escherichia coli ATCC accession No. 25922, and Staph. aureus ATCC accession No. 13301. Generally, the target/test bacteria are cultured in vitro in Mueller-Hinton broth (BBL Microbiology Systems, Casksonville, 199).
- 35 broth (BBL Microbiology Systems, Cockeysville, MD) for 18 hours at 37°C.

Typically, cultures of a test pathogen will be prepared by suspending colonies grown on solid medium (for example, trypticase soy agar plates) into 70 ml of Mueller-Hinton broth so that a culture with an optical density of about 0.1 at 540 nm results. Appropriate dilutions of the bacterial cells are then prepared in DPBS.

Animals. Typically, any acceptable animal model may be used to assess the efficacy of the antibacterial oligonucleotides. Additionally, experimental protocols and 10 conditions will necessarily be adjusted as applicable depending on the bacterial pathogen being tested and the mode of infection. Accordingly, the following example is provided merely for purposes of exemplification and should not be deemed as limiting the present invention in any way 15 whatsoever.

Six- to eight-week-old CD1 mice or NMRI mice, 24-28 g in size, are typically used in these studies. The CD1 strain of mouse has been used in the past for certain studies of infectious diseases and therapeutics (e.g., Brogden et al., (1986); Cavalieri et al., (1991); Lister and Sanders, Antimicrob. Agents Chemother. 39:930-936 (1995)), as has the NMRI strain (Hof et al., Infection 114:190-194 (1986)).

Thus, both of the above strains are exemplary of well established infectious disease models that are also readily available to those of ordinary skill.

Typical animal tests comprise a minimum of about 5-8 animals in each treatment group (1 cage of 5 mice each) in order to demonstrate adequately the statistical reproducibility of a given experimental observation. By 30 using at least about 5 test animals, one can compensate for variabilities such as differing growth rates of microorganisms in a given animal and any variables introduced by the repeated handling and injection of the animals.

Injection of microorganisms. Test animals are typically 35 injected subcutaneously (SC) on the back (intrascapular) with approximately 0.3 ml of bacterial cell suspension in 1.5% liquified sterile tryptose phosphate agar held at 39°C

essentially as described by Hof et al. (1986) or I.P. with 5% mucin (Lister & Sanders, 1995).

Administration of Oligonucleotides. At the time of injection of bacteria or at various times after injection 5 with the indicated microorganism, the test animals are treated by administration of a bolus injection of oligonucleotides at, for example, 0, 1.0, 2.5, 5.0 or 10.0 mg/kg (5 separate groups, one dose per group of 12 animals) to determine optimum therapeutic dose of a given

- 10 antibacterial oligonucleotide. The oligonucleotide is generally administered I.P. in a volume of approximately 0.5 ml of sterile saline, using a sterile 25-gauge needle or through an Alzets pump. Optionally, the solution comprising the antibacterial oligonucleotide may also be administered
- 15 I.V., subcutaneously, orally, or by any other means suitable for the given pathogen being tested.

Where applicable, bacteremia will be monitored by collecting daily blood samples from two animals from each group. One fully-anesthetized animal from the negative 20 control group (no bacterial infection) will be bled by cardiac puncture and subsequently euthanized. The number of colony forming units (CFU) in the blood samples will then be determined by plating samples on agar and doing bacterial

The minimum lethal dose for a given bacterial pathogen, e.g., Escherichia coli ATCC accession No. 25922, is determined for CFI mice after the pathogen is injected I.P. in 0.5 ml DPBS or S.C. plus agar. The minimum lethal inoculum is the minimum dose that results in the death of all 30 of the test subjects during the five to seven days postinfection.

colony assays.

Alternatively, female NMRJ mice may be used with, for example, Escherichia coli ATCC accession No. 25922, which is known to cause animal death within five to seven days after 35 intra-clavicular injection.

The dose of antibacterial oligonucleotide that protects 50% of the test animals from death (protective doses $50\%-PD_{50}$)

is determined as follows. Beginning at various times after injection of the bacterium into the test animals, and continuing for four days thereafter, the antibacterial oligonucleotide (or its control) is injected S.C. into the 5 test animals in about 0.15 ml DPBS at final concentrations that will vary as appropriate for the given assay. For example, about 0.0, 1.0, 2.0, 2.5, and 5.0 mg/kg of antibacterial oligonucleotide may typically be used. Animals surviving for more than five to seven days after initial 10 bacterial inoculation will be maintained an additional seven days, and then euthanized by CO, asphyxiation for further study. Optionally, the test animals are maintained for more extended periods after initial infection in order to assess the long-term efficacy of oligonucleotide treatment.

A similar bacterial inoculation and oligonucleotide treatment protocol can be used to determine the kinetics of bacteria clearance from the peripheral blood of bacteremic animals after treatment with antibacterial oligonucleotide. In these studies, groups of twelve animals each are infected as above with Escherichia coli, and a group of six mice is sham injected with only saline (the control group). The groups of infected mice are then treated with (a) saline or (b) oligonucleotide, while the control group is only treated with saline. At suitable time periods post-infection, blood samples are taken, and the number of test pathogen cells per ml of blood is determined by standard dilution and culture methods.

The above animal models are merely exemplary of the myriad of animal models that may be used to establish the 30 efficacy of the presently described antibacterial oligonucleotides, and many other modalities for testing the claimed invention are available to one of ordinary skill. For example, the LD₅₀ of a given pathogen may be established (or previously known), and the efficacy of the antibacterial oligonucleotide determined, testing whether substantially all of the test animals survive bacterial exposure.

Additionally, immunocompromised animals may also be used, *i.e.*, nude mice, SCID mice, etc., to study the antibacterial effects of the described oligonucleotides in the absence of a correctly functioning immune system.

5

4.7. Pharmaceutical Compositions And Delivery

Pharmaceutical compositions containing the oligonucleotides of the invention in intimate admixture with a pharmaceutical carrier can be prepared according to 10 conventional pharmaceutical compounding techniques. The carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g., intravenous, oral, topical, aerosol (for topical or inhalation therapy), suppository, parenteral, or spinal 15 injection.

In preparing the compositions in oral dosage form, any of the usual pharmaceutical media may be employed, such as, for example, water, glycols, oils, alcohols, flavoring agents, preservatives, coloring agents and the like in the case of oral liquid preparations (such as, for example, suspensions, elixirs, and solutions); or carriers such as starches, sugars, diluents, granulating agents, lubricants, binders, disintegrating agents and the like in the case of oral solid preparations (such as, for example, powders,

- 25 capsules and tablets). Because of their ease in administration, tablets and capsules represent the most advantageous oral dosage unit form, in which case solid pharmaceutical carriers are obviously employed. If desired, tablets may be sugar-coated or enteric-coated by standard
- 30 techniques. Oral dosage forms of antibacterial oligonucleotides will be particularly useful for the treatment of bacterial infections of the gastrointestinal tract and ulcers caused by or associated with bacterial infection (e.g., Helicobacter pylori infection, and the
- 35 like). Additionally, given that bacterial infection has been associated with hyperproliferative disorders of the immune system (i.e. inflammatory bowel disease), the presently

described antibacterial oligonucleotides may be used to treat hyperproliferative disorders including, but not limited to, Crohn's disease and ulcerative colitis by specifically eliminating the causative or contributory microorganisms from the bacterial flora of the gut.

For parenteral application by injection, preparations may comprise an aqueous solution of a water soluble, or solubilized, and pharmaceutically acceptable form of the antibacterial oligonucleotide in an appropriately buffered 10 saline solution. Injectable suspensions may also be prepared using appropriate liquid carriers, suspending agents, pH adjusting agents, isotonicity adjusting agents, preserving agents, and the like may be employed. Actual methods for preparing parenterally administrable compositions and 15 adjustments necessary for administration to subjects will be known or apparent to those skilled in the art and are described in more detail in, for example, Remington's Pharmaceutical Science, 15th Ed., Mack Publishing Company, Easton, Pa (1980), which is incorporated herein by reference. 20 The presently described oligonucleotides should be parenterally administered at concentrations below the maximal tolerable dose (MTD) established for the antibacterial

For topical administration, the carrier may take a wide 25 variety of forms depending on the preparation, which may be a cream, dressing, gel, lotion, ointment, or liquid.

oligonucleotide.

Aerosols are prepared by dissolving or suspending the oligonucleotide in a propellant such as ethyl alcohol or in propellant and solvent phases. The pharmaceutical 30 compositions for topical or aerosol form will generally contain from about 0.01% by weight (of the oligonucleotide) to about 40% by weight, preferably about 0.02% to about 10% by weight, and more preferably about 0.05% to about 5% by

Suppositories are prepared by mixing the oligonucleotide with a lipid vehicle such as theobroma oil, cacao butter, glycerin, gelatin, or polyoxyethylene glycols.

weight depending on the particular form employed.

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The presently described antibacterial oligonucleotides may be administered to the body by virtually any means used to administer conventional antibiotics. A variety of delivery systems are well known in the art for delivering 5 bioactive compounds to bacteria in an animal. These systems include, but are not limited to, intravenous or intramuscular or intrathecal injection, nasal spray, aerosols for inhalation, and oral or suppository administration. The specific delivery system used depends on the location of the 10 bacteria, and it is well within the skill of one in the art to determine the location of the bacteria and to select an appropriate delivery system.

The present invention is further illustrated by the following examples, which are not intended to be limiting in 15 any way whatsoever.

5.0. EXAMPLES

5.1. Oligonucleotide Synthesis

Oligonucleotides were synthesized using commercial 20 phosphoramidites on commercially purchased DNA synthesizers at either 1 μ M or 15 μ M scales using standard phosphoramidite chemistry. Oligonucleotides were deprotected following phosphoramidite manufacturers protocols. Oligonucleotides to be used unpurified were either dried down under vacuum or 25 precipitated and then dried.

Sodium salts of oligonucleotides were prepared using the commercially available DNA-Mate (Barrskogen, Inc.) reagents or conventional techniques such as the commercially available exchange resin, e.g., Dowex (Tradename), or by addition of sodium salts followed by precipitation, diafiltration, or gel filtration, etc.

Oligonucleotide preparations that would be subject to further purification were initially chromatographed on commercially available reverse phase or ion exchange media

35 (preferably, SAX, strong anion exchange media) such as Source Q made by Pharmacia, Toyopearl super Q made by Tosohaas, Protein Pak made by Waters, Macroprep Q made by BioRad, and

the like. Peak fractions were combined and the samples desalted and concentrated by ethanol precipitation, diafiltration, or gel filtration followed by lyophilization or solvent evaporation under vacuum in commercially available instrumentation such as Savant's Speed Vac. Optionally, the oligonucleotides may also be electrophoretically purified using polyacrylamide gels.

A variety of commercially available gel filtration media are particularly well suited for the desalting and/or

10 purification of antibacterial oligonucleotides. Gel filtration media which may be used include Sephadex or Superdex made by Pharmacia, Trisacryl made by BioSepra, BioGel (preferably P-series, or more preferably P4) made by BioRad, Toyopearl HW SEC made by Tosohaas, Cellufine made by Amicon, and the like. Optionally, the gel filtration step may be repeated several times in order to better remove low molecular weight species, and particularly alkyl amines and/or alkyl ammonium compounds, from the oligonucleotide preparations.

- Cation exchange columns comprising media such as Macroprep S (or CM) made by BioRad (preferably in the NH₄: form), Dowex resins, or Amberlite resins are also useful to remove contaminants from antibacterial oligonucleotide preparations. Typically, the pH of the eluted

 25 oligonucleotide will be increased to about 7-8 using ammoniants
- 25 oligonucleotide will be increased to about 7-8 using ammonium hydroxide consequential to this step.

Alternatively, exhaustive dialysis or diafiltration may be used to remove salts or contaminants that inhibit or mask the antibacterial activity of the oligonucleotides (e.g.,

- 30 alkyl amines and/or alkyl ammonium compounds). Exhaustive butanol extractions, chloroform extraction followed by ethanol washes or multiple ethanol extractions may be used to obtain purified oligonucleotides that retain antibacterial activity.
- Oligonucleotides to be used in bacterial experiments were dissolved in pyrogen free, sterile, physiological saline (i.e., 0.85% saline), sterile Sigma H₂O, and filtered through

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a 0.45 micron Gelman filter (or a sterile 0.2 micron pyrogen free filter prior to animal studies). Table 1 contains a list of all oligonucleotide sequences used in the examples. Although the majority of oligonucleotides used in the

- 5 examples were constructed using a phosphorothicate backbone, unless otherwise noted, it should be understood that any of a wide variety of chemical backbones could be also used to generate oligonucleotides comprising the sequences listed in Table 1. The antibacterial oligonucleotides were tested
- 10 for inhibition (INH) activity against drug resistant Gram negative (Escherichia coli ATCC accession No. 35218) and Gram positive (Staphylococcus aureus ATCC accession No. 13301) microorganisms. The percent inhibition data in Table 1 were averaged and normalized to a concentration of 2 mg/ml.
- Tables 2(A-W) provide time course experiments that test the inhibitory activity (against Escherichia coli ATCC accession No. 35218 or Staphylococcus aureus ATCC accession No. 13301) of the indicated oligonucleotides when present at 2 mg/ml in the culture medium as targeted against genes that
- 20 represent nearly all known gene classes in bacteria. In brief, Table 2A shows the inhibitory effect of oligonucleotide 28 (NBT 28, SEQ ID NO. 1); Table 2B tests oligonucleotide 10 (SEQ ID NO. 17); Table 2C tests oligonucleotide 43 (SEQ ID NO. 34), Table 2D shows the
- 25 inhibitory effect of oligonucleotide 27 (SEQ ID NO. 45); Table 2E tests oligonucleotide 2 (SEQ ID NO. 120); Table 2F tests oligonucleotide 89 (SEQ ID NO. 61); Table 2G tests oligonucleotide 103 (SEQ ID NO. 64); Table 2H tests oligonucleotide 132 (SEQ ID NO. 65), Table 2I shows the
- 30 inhibitory effect of oligonucleotide 19 (SEQ ID NO. 66);
 Table 2J tests oligonucleotide 16 (SEQ ID NO. 72); Table 2K tests oligonucleotide 96 (SEQ ID NO. 79); Table 2L tests oligonucleotide 21 (SEQ ID NO. 85); Table 2M shows the inhibitory effect of oligonucleotide 18 (SEQ ID NO. 95);
- 35 Table 2N tests oligonucleotide 105 (SEQ ID NO. 103); Table 20 tests oligonucleotide 46 (SEQ ID NO. 105); Table 2P tests oligonucleotide 114 (SEQ ID NO. 112); Table 2Q tests

oligonucleotide 32 (SEQ ID NO. 116); Table 2R tests oligonucleotide 73 (SEQ ID NO. 124); Table 2S tests oligonucleotide 63 (SEQ ID NO. 130), Table 2T shows the inhibitory effect of oligonucleotide 78 (SEQ ID NO. 134); Table 2U tests oligonucleotide 71 (SEQ ID NO. 151); Table 2V tests oligonucleotide 14 (SEQ ID NO. 154); and Table 2W tests oligonucleotide 5 (SEQ ID NO. 152).

5.2. MIC With Escherichia coli

Oligonucleotides from <u>every</u> known gene class in bacteria were used to test inhibition of bacterial growth in a modified MIC test (described above). In all cases the control bacterial cells entered exponential growth while the test cells to which oligonucleotide had been added showed no growth at all or significant inhibition of growth (see Table 1).

Similar results were achieved with other oligonucleotides selected using the parameters described above, which were subsequently synthesized, purified and 20 tested using the same MIC analysis. See Table 1.

The results in Table 1 demonstrate that antisense or antigene (inhibition of expression by DNA triplex formation) oligonucleotides are effective against a variety of genes. For example: genes involved in energy metabolism (sugar 25 metabolism, fatty acid metabolism), cell division (DNA replication, cell wall biosynthesis), global regulatory proteins, protein synthesis (tRNA synthesis, mRNA stability, rRNA synthesis, ribosomal protein, translation factors), virulence factors, cell wall and membrane synthesis (fatty

- 30 acid and phospholipid synthesis, lipopolysaccharide synthesis, periplasmic-secretory proteins, transport proteins, outer-membrane proteins), amino acid biosynthesis, nucleic acid synthesis, nitrate reductase, vitamin metabolism, and drug resistance.
- In fact, Figure 2 shows that the described antibacterial oligonucleotides proved effective against a wide variety of genes from both Gram negative and Gram positive bacteria.

More specifically, oligonucleotides targeted against bacterial genes relating to: energy metabolism (A); DNA replication (B); cell division (C); regulatory proteins (D); cell wall biosynthesis (E); sugar metabolism (F); virulence, 5 pili, flagella (G); fatty acid metabolism (H); mRNA synthesis (I); tRNA synthesis (J); rRNA synthesis (K); ribosomal protein synthesis (L); protein synthesis (M); phospholipid synthesis (N); periplasmic/secretory protein synthesis (O); regulation and synthesis of transport proteins (P); amino 10 acid biosynthesis and metabolism (Q); lipopolysaccharide synthesis (R); purine/pyrimidine biosynthesis and metabolism (S); outer membrane protein synthesis and regulation (T); nitrate reductase synthesis and regulation (U); drug resistance (V); and vitamin metabolism and biosynthesis (W) 15 were capable of significantly inhibiting the growth of both Gram negative and Gram positive bacteria.

Thus, antibacterial oligonucleotides were effective against virtually every major cellular function tested (as determined by the MIC assay).

- As additional genome sequence data are obtained for bacteria, this invention may be extended to oligonucleotide targets within newly described bacterial sequences.

 Antibacterial oligonucleotides may be constructed with a range of backbones including, but not limited to:
- 25 phosphorothioates; p-ethoxy oligonucleotides (partially or fully substituted); or 2'-O-methyl oligonucleotides (partially or fully substituted). Oligonucleotides comprising all of the above backbones have proved equally effective in inhibiting bacterial growth. In view of the
- 30 effectiveness of oligonucleotides comprising the chemical backbones listed above, chimeric oligonucleotides (comprising mixed backbones) are also deemed to be effective antibacterial agents.

Several oligonucleotides based on the NBT 18 sequence 35 (SEQ ID NO. 95) were also capable of inhibiting the growth of two clinically relevant pathogens that have proven resistant to most conventional antibiotics - Escherichia coli clinical

isolate ATCC accession No. 35218 (Tables 3A and 3B), and Staphylococcus aureus clinical isolate ATCC accession No. 13301 (Tables 3C and 3D). The NBT 18 sequence variations that were tested in Tables 3A and 3B include: A - the NBT 18 5 sequence with a 2'-O-Methoxy substituted backbone; B - a truncated (12mer, SEQ ID NO. 174) version of the NBT 18 sequence with a phosphorothioate backbone; C - a truncated (15mer, SEQ ID NO. 175) region of the NBT 18 sequence with a phosphorothioate backbone; D - a truncated (15mer) region of 10 the NBT 18 sequence with a phosphorothicate backbone and a 5' amino group; and E - the NBT 18 sequence with a phosphorothioate backbone. The NBT 18 sequence variations that were tested in Tables 3C and 3D include: A - the NBT 18 sequence with a 2'-O-Methoxy substituted backbone; B - the 15 NBT 18 sequence with a p-ethoxy substituted backbone; C - a truncated (12mer) region of the NBT 18 sequence with a phosphorothicate backbone; D - a truncated (15mer) region of the NBT 18 sequence with a phosphorothicate backbone; and E a truncated (18mer, SEQ ID NO. 176) region of the NBT 18 20 sequence with a phosphorothicate backbone. Tables 3(A-D) indicate that the observed antibacterial effect was largely a feature of the antisense sequence of NBT 18 instead of the backbone of a given oligonucleotide (i.e., nonspecific sulphur effects, etc.).

These data further indicate that oligonucleotides comprising less than one half of the full-length (27 base) sequence of NBT 18 retain the ability to inhibit the growth of at least two clinically significant pathogens.

5.3. MIC With Gram Negative And Gram Positive Bacteria A representative number of the antisense

oligonucleotides were tested against a wide variety of bacterial species including Streptococcus (Streptococcus mutans (ATCC accession No. 25175)), Streptococcus pyogenes

35 (ATCC accession No. 14289), Streptococcus pneumoniae or Pneumococcus pneumoniae (ATCC accession No. 39937), and Streptococcus faecalis or Enterococcus faecalis (ATCC

accession No. 19433), Staphylococcus aureus (ATCC accession No. 29213), Staphylococcus aureus (ATCC accession No. 13301), Escherichia coli (ATCC accession Nos. 11370, 25922, and 29214), Salmonella typhimurium (ATCC accession No. 23564),

- 5 Pseudomonas fluorescens (ATCC accession No. 13525),
 Klebsiella pneumoniae (ATCC accession No. 4352), Serratia
 liquefaciens (ATCC accession No. 27592), Neisseria sicca
 (ATCC accession No. 9913), Mycobacterium smegmatis (ATCC
 accession No. 19420), Yersinia mollareti (ATCC accession No.
- 10 43969), Haemophilus segnis (ATCC accession No. 33393), Haemophilus vaginalis (ATCC accession No. 14018), Shigella sp. (ATCC accession No. 11126), Vibrio fischeri (ATCC accession No. 7744), and Helicobacter mustelae (ATCC accession No. 43772).
- Representative data generated with phosphorothicate forms of the oligonucleotides are provided in Tables 4(A-Z). In brief, antibacterial oligonucleotides nos. 18 (SEQ ID NO. 73), 39 (SEQ ID NO. 30), 63 (SEQ ID NO. 130), 78 (SEQ ID NO. 134), and 73 (SEQ ID NO. 124) were tested against Salmonella
- 20 typhimurium (Tables 4A and 4B); antibacterial
 oligonucleotides 39 (SEQ ID NO. 30), 63 (SEQ ID NO. 130), 78
 (SEQ ID NO. 134), 82 (SEQ ID NO. 161), and 114 (SEQ ID NO.
 112) were tested against Pseudomonas aeruginosa (Tables 4C
 and 4D); antibacterial oligonucleotides 114 (SEQ ID NO. 112),
- 25 78 (SEQ ID NO. 134), 73 (SEQ ID NO. 124), 71 (SEQ ID NO. 151), and 111 (SEQ ID NO. 132) were tested against Klebsiella pneumoniae (Tables 4E and 4F); antibacterial oligonucleotides 2 (SEQ ID NO. 50), 4 (SEQ ID NO. 173), 127 (SEQ ID NO. 143), 63 (SEQ ID NO. 130), and 73 (SEQ ID NO. 124) were tested
- 30 against Yersinia mollaretti (Tables 4G and 4H); antibacterial oligonucleotides 16 (SEQ ID NO. 72), 12 (SEQ ID NO. 80), 20 (SEQ ID NO. 84), 3 (SEQ ID NO. 121), and 15 (SEQ ID NO. 81) were tested against Neisseria sicca (Tables 4I and 4J); antibacterial oligonucleotides 2 (SEQ ID NO. 50), 39 (SEQ ID
- 35 NO. 30), 82 (SEQ ID NO. 161), and 114 (SEQ ID NO. 112) were tested against Serratia liquefaciens (Table 4K); antibacterial oligonucleotides 1 (SEQ ID NO. 119), 89 (SEQ ID

NO. 61), 127 (SEQ ID NO. 143), 132 (SEQ ID NO. 15), and 114 (SEQ ID NO. 112) were tested against Streptococcus mutans (Tables 4L and 4M); antibacterial oligonucleotides 1 (SEQ ID NO. 119), 89 (SEQ ID NO. 61), 127 (SEQ ID NO. 143), 132 (SEQ 5 ID NO. 15), and 114 (SEQ ID NO. 112) were tested against Streptococcus pyogenes (Tables 4N and 40); antibacterial oligonucleotides 1 (SEQ ID NO. 119), 89 (SEQ ID NO. 61), 127 (SEQ ID NO. 143), 132 (SEQ ID NO. 15), and 114 (SEQ ID NO. 112) were tested against Shigella (Tables 4P and 4Q); 10 antibacterial oligonucleotide 78 (SEQ ID NO. 134) was tested against Haemophilus (Table 4R); antibacterial oligonucleotides 114 (SEQ ID NO. 112), 10 (SEQ ID NO. 17), 21 (SEQ ID NO. 85), 18 (SEQ ID NO. 73), and 78 (SEQ ID NO. 134) were tested against Mycobacterium (Tables 4S and 4T); 15 antibacterial oligonucleotide 78 (SEQ ID NO. 134) was tested against Helicobacter (Table 4U); antibacterial oligonucleotides 89 (SEQ ID NO. 61), 127 (SEQ ID NO. 143), 132 (SEQ ID NO. 15), p127 (SEQ ID NO. 143 with a p-Ethoxy backbone), 1 (SEQ ID NO. 119), and 76 (SEQ ID NO. 127) were 20 tested against Enterococcus (Tables 4V and 4W); antibacterial oligonucleotides 1 (SEQ ID NO. 119), 78 (SEQ ID NO. 134), 114 (SEQ ID NO. 112), 127 (SEQ ID NO. 143), and 132 (SEQ ID NO. 15) were tested against Streptococcus pneumonia (Tables 4X and 4Y); and antibacterial oligonucleotides 78 (SEQ ID NO. 25 134) and 127 (SEQ ID NO. 143) were tested against Vibrio (Table 4Z). The data in Tables 4A-Z indicate that the antibacterial oligonucleotides targeted to varying classes of genes are capable of strongly inhibiting the growth of a broad spectrum of bacterial species. No significant 30 difference in antibacterial activity was found when different

Additionally, Figures 3(a-c) respectively provide time course data providing percent inhibition as a function of 35 time for oligonucleotides 73 (SEQ ID NO. 124), 63 (SEQ ID NO. 130), and 18 (SEQ ID NO. 73) as measured against Salmonella typhimurium; Figures 4(a-c) respectively provide time course

stereoisomers of phosphorothicate backbone oligonucleotides

were tested.

data showing percent inhibition as a function of time for oligonucleotides 39 (SEQ ID NO. 30), 78 (SEQ ID NO. 134), and 63 (SEQ ID NO. 130) as measured against *Pseudomonas* aeruginosa; and Figures 5(a-b) respectively provide time 5 course data showing percent inhibition as a function of time for oligonucleotides 73 (SEQ ID NO. 124) and 114 (SEQ ID NO. 112) as measured against *Klebsiella pneumoniae*.

In view of the wide range of bacteria already successfully tested, any oligonucleotides chosen and prepared 10 in the manner described herein will be equally effective against a given bacterial target. In addition to the species explicitly mentioned herein, a wide variety of other bacterial pathogens may be treated using the described compositions. A relatively comprehensive review of such 15 pathogens is provided, inter alia, in Mandell et al., 1990, Principles and Practice of Infectious Disease 3rd. ed., Churchill Livingstone Inc., New York, N.Y. 10036, herein incorporated by reference.

20 5.4. MIC At 24 Hours

In order to distinguish whether the antibacterial oligonucleotides had transient bacteriostatic effects, or long lasting effects, MIC assays were extended to include a time point of over 24 hours. These data are presented in

- 25 Tables 5A-D. Tables 5A and 5B show, inter alia, that oligonucleotides 21 (SEQ ID NO. 156), 68 (SEQ ID NO. 148), and 85 (SEQ ID NO. 106), 112 (SEQ ID NO. 62), and 18 (SEQ ID NO. 73) continue to substantially inhibit the growth of Staphylococcus aureus ATCC accession No. 13301, for at least
- 30 25 hours. These data indicate that the tested oligonucleotides have long-term bacteriostatic or bactericidal (see Figure 9, below) effects on Staphylococcus aureus ATCC accession No. 13301. Moreover, the timing of antibacterial oligonucleotide addition does not significantly
- 35 affect the observed antibacterial activity since activity was seen when the addition of antibacterial oligonucleotide was delayed for 180, 350, or 480 min.

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Conversely, Tables.5C-D indicates that, although a substantial amount of growth inhibition occurs initially, the same oligonucleotides do not significantly inhibit the growth of Escherichia coli ATCC accession No. 35218 when growth was 5 assayed 27 hours after the bacteria were initially exposed to the oligonucleotides. The data in Tables 5C and 5D indicate that oligonucleotides 21 (SEQ ID NO. 156), 68 (SEQ ID NO. 148), 85 (SEQ ID NO. 106), 112 (SEQ ID NO. 62), and 18 (SEQ ID NO. 73) are bacteriostatic for Escherichia coli ATCC 10 accession No. 35218. Escherichia coli ATCC accession No. 35218 represents a particularly virulent, multiple drug resistant strain of Escherichia coli. When oligonucleotide number 89 (SEQ ID NO. 61) was tested against Escherichia coli accession No. 25922, a moderately penicillin resistant 15 strain, a dose-dependent long lasting bacteriostatic effect was observed (see Tables 5E and 5F). It is expected that multiple doses of the same oligonucleotide, rather than a single dose, might result in enhanced long-term activity against the more resistant Escherichia coli ATCC accession 20 No. 35218.

The 24-hour MIC studies were performed essentially as described above with the exceptions that: growth of the target bacteria to reach an OD₅₂₅ of 0.1 occurs in approximately 8 hours instead of about 12 to 16 hours; 25 bacterial growth is monitored throughout the experiment as well as at the end-points; and an additional test was conducted that used starved cells as the initial inoculum instead of fresh log cultures (which provided similar antibacterial results).

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5.5. Purification Studies

The MIC test was carried out as described in Section 4.5., supra. The test oligonucleotides received various post-synthesis treatments, and the percent inhibition of the cell culture growth was calculated as described supra. See Tables 6A and 6B.

Oligonucleotide NBT. 78 (SEQ ID NO. 134), was given the following treatments:

- A. butanol precipitated and resuspended as an ammonium salt;
- 5 B. butanol precipitated, converted to a sodium salt, desalted on a gel filtration column (described Section 5.1);
 - C. purified via anion exchange HPLC, desalted by gel filtration;
- D. butanol precipitated, converted to a sodium salt, desalted on a reverse phase HPLC column (trityl off);

15

25

- E. butanol precipitated, ammonium hydroxide added, desalted via gel filtration, left as an ammonium salt;
 - F. butanol precipitated <u>once</u>, filtered through a 0.45 micron filter (e.g., Gelman Acrodisc, Millipore, Nalgene, etc.) followed by ethanol precipitation;
- G. butanol precipitated <u>twice</u>, filtered through a 0.45
 micron filter (e.g., Gelman Acrodisc, Millipore,
 Nalgene, etc.), and washed three times with 95%
 ethanol;
 - H. butanol precipitated <u>twice</u>, filtered through a 0.45 micron filter (e.g., Gelman Acrodisc, Millipore, Nalgene, etc.), washed with chloroform and ethanol;
 - I. butanol precipitated <u>twice</u>, filtered through a 0.45 micron filter (e.g., Gelman Acrodisc, Millipore, Nalgene, etc.), butanol precipitated 2 more times, and washed once with ethanol.
- The results in Tables 6A and 6B demonstrate that the protocol used to purify the oligonucleotides greatly affects bacterial susceptibility in a MIC test. Oligonucleotides that are treated only by butanol precipitation inhibited bacterial growth by less than 25 percent. However,
- 35 oligonucleotides that were subject to: a) gel filtration; b) four butanol precipitations; or c) two butanol extractions, followed by ethanol or chloroform extractions all

demonstrated greater than 85% inhibition of the growth of the test bacteria used in the MIC assay (see B, C, E, G, H and I). Oligonucleotides may also be purified by strong anion exchange (SAX) chromatography, reverse-phase chromatography, 5 strong cation exchange (SCX) chromatography, followed by size exclusion chromatography (SEC). Alternatively, after the first SCX column, a second SCX column can be run followed by a reverse-phase chromatography step. Optionally, the SCX step may be supplemented or replaced by an alcohol (e.g., 10 ethanol, etc.) precipitation step.

The above results demonstrated that proper post synthesis handling protocols play an integral role in the production of oligonucleotides that display antibacterial activity.

15 There are a variety of contaminants that may be present in an oligonucleotide preparation after cleavage from the solid supports and removal of the protecting groups, and even after HPLC treatment. These contaminants include residual protecting groups, and contaminants that are introduced or 20 generated during synthesis or purification. Examples of such contaminant include, but are not limited to, quaternary amines (particularly alkyl amines and/or alkyl ammonium compounds), acetamide, acetic acid, 2-cyanoethanol, isobutyramide, isobutyric acid, benzamide, benzoic acid, 25 succinimide, succinic acid, t-butylphenoxyacetamide (or acetic acid), phenoxyacetamide (or acetic acid). Given the results shown in Tables 6A and 6B, it is clear that the substantial removal of the above or other contaminants greatly enhances the antibacterial activity of an 30 oligonucleotide.

Contaminants that are particularly important to remove from the oligonucleotide preparations include compounds that directly or indirectly inhibit bacterial uptake of the oligonucleotides, or otherwise mask the antibacterial effects of the oligonucleotides. One way that a contaminant may mask the antibacterial efficacy of an oligonucleotide is by stimulating bacterial growth in a manner that effectively

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compensates for the antibacterial activity of a given oligonucleotide. Accordingly, the present finding that certain contaminants (i.e., alkyl amines and/or alkyl ammonium compounds) that are typically present in 5 conventional oligonucleotide preparations may mask the in vitro antibacterial activity of oligonucleotides represents a seminal discovery that requires a fundamental reassessment of

In particular, an impurity in anion exchange (AX) HPLCpurified modified linkage oligonucleotides has been isolated
and partially characterized which stimulates bacterial growth
both in vitro and in vivo. This impurity/stimulatory
material is a mixture of small, polar, multialkyl amino or
15 alkyl ammonium compounds that have negligible absorbance at
254 nm. The impurity is apparently generated from the AXHPLC stationary phase during the elution gradient.

the utility of oligonucleotides as antibacterial agents in

vivo.

The absence of an active chromophore at 254 nm effectively renders the impurity invisible to the absorbance 20 detectors used during HPLC of DNA oligonucleotides. Since anion exchange chromatography precludes the use of conductivity detectors to monitor peaks, the impurity is also virtually invisible during the purification and analytical HPLC procedures typically used in the manufacture of 25 oligonucleotides.

As shown above, the impurity can be removed and isolated from the oligonucleotide preparations by using a series of desalting steps. For example, in the first step, the oligonucleotide was concentrated by first loading the pooled 30 fractions of an AX purification run onto appropriately sized Hamilton PRP-1 or PRP-3 columns. The salt was then removed from the column by washing with water until the conductivity of the wash eluant was below 25 μ S/cm. Finally, the oligonucleotide was eluted as a concentrated solution (app. 35 100-300 OD's per mL) using a moderately steep (5% per minute) gradient of water:90% ethanol. It should also be noted that oligonucleotides purified in this manner must contain at

least two phosphorothicate or p-ethoxy linkages, or some other non-polar modification in order to adequately absorb to the stationary phase.

In the second step, the oligonucleotide solution was 5 concentrated or removed entirely by lyophilization prior to further purification by size-exclusion chromatography (SEC). The oligonucleotide was re-suspended in a minimum amount of water prior to application to the SEC column. Since essentially all of the salt from the AX purification was 10 removed by the RP step, the oligonucleotide was dissolved in a relatively small volume of water. This small volume helps maximize resolution in the SEC step.

A column was prepared using virgin BioGel P-4 medium or fine particle SEC medium, using a modified manufacturer's 15 procedure to swell the medium. The column used was 45-50 cm long and 2.2 cm diameter. The flow rate was approximately 1-2 mL/minute. This size column can be used to purify 1,000-3,000 OD's of modified linkage oligonucleotides that are at least 12 bases in length. If the oligonucleotides have more than 30% phosphorothicate linkages, the maximum loading drops to about 2,000 OD's. Columns and sample sizes may be scaled up as long as a flow velocity of about 30-75 cm/hr is maintained, and the column height remains at least about 40 cm.

The oligonucleotides were eluted with water while monitoring the conductivity and the absorbance at 254 nm. The purification may be easily be modified by monitoring at 280 nm, and the like. Collection began when the oligonucleotide concentration became appreciable (as measured 30 by 0.D.), and stopped at no later than about 8 minutes after collection began. If, after the conductivity initially rose, it fell and then began to rise again, collection was terminated. It was important to stop collection as described because oligonucleotides collected after this point typically included the stimulatory impurities.

The collected oligonucleotide solutions were checked for concentration and lyophilized. Typically, the above protocol

agreement beginner dem fact to the control of the c

resulted in the purified oligonucleotides having the desired antimicrobial activities.

When separation continued after the collection of the oligonucleotide peak, several other peaks were seen which 5 displayed little to no absorbance at 254 nm, but noticeable conductivity. The amount of impurity observed varied for each individual purification. The variation was probably attributable to the different salt concentrations required to elute different oligonucleotides, or variations in the length 10 of time since the AX column was last used, etc.

While the detected amounts of impurity generally remained a small percentage of the net composition, both in vivo and in vitro testing showed that the impurities stimulate bacterial growth. Oligonucleotides that were not purified by AX-HPLC but are otherwise treated the same did not display either of the peaks observed during SEC, and did not have a stimulatory effect. However, oligonucleotides that were AX-HPLC purified and desalted as described, but were not further purified by SEC showed either stimulatory effects or, where the amounts of the impurities were not high, neutral or a significantly reduced antibiotic effect.

Spectroscopic analysis (1H -NMR, A_{254} absorbance, GC-MS, and FAB and ESI positive ion mass spectrometry) pointed to a comparatively small, simple molecule, or mixture of similar

- 25 components, that were eluted along with the oligo. These compound(s) coeluted with oligonucleotide during the reversephase concentration/desalting process. In particular, analysis by electrospray mass spectroscopy of small molecular weight material removed from an oligonucleotide preparation
- 30 that had been purified on a Waters Protein Pak 40Q revealed complex mixture of amino compounds with the common feature of signals at m/z 58 and m/z 72. These two signals are derived from the N,N-diethyl-N-(2-hydroxypropyl) quarternary amino functional group used as the cationic absorption moiety on
- 35 the Protein Pak Q SAX stationary phase. Electrospray analysis of similar material from a N,N,N-trimethyl quarternary amino polymer-based SAX phase (e.g., BioRad's

Macroprep Q) also yielded equivalent signals indicative of the cleavage of absorption sites from the stationary phase. These low molecular weight materials were removed by SEC, and were also removed by a combination of SCX and reverse phase 5 chromatography.

The steep ramping required for concentration purposes did not permit conditions suitable for resolution of close-running materials. However, the SEC step outlined above was capable of sufficiently removing the impurities to allow the 10 detection of a consistent pattern of antibiotic activity inherent in the presently described purified oligonucleotides. Accordingly, the SEC step provides a process that allowed for the consistent and predictable removal of the stimulatory impurities from the 15 oligonucleotide preparations.

As discussed above, oligonucleotides that have been purified using different procedures (i.e., no chromatography steps) consistently showed antibiotic effects that were comparable to the oligonucleotides purified as outlined 20 immediately above.

In some very non-polar oligonucleotides, such as total p-ethoxy and chimeras with p-ethoxy/2'-O-methyl RNAs components, the concentration of ethanol required to elute the oligonucleotides from the reverse-phase column was high enough to allow some removal of the low-absorbing high conductivity material prior to the elution of the oligonucleotides. However, the resolution was not sufficiently clean to allow straight-forward characterization. This separation was not observed with predominantly S-oligonucleotides.

The ability of the RP-column to provide any separation may also be affected by the base composition of the oligonucleotides as well as the type of linkages employed to construct the oligonucleotides. Typically, the use of 35 ethanol provided more control over the elution process than acetonitrile, which has higher elution power than ethanol.

Additionally, the use of ethanol during this step has implications for cGMP validation.

Another feature of the RP step is that the great reduction of inorganic salt during the reverse-phase protocol allows for the use of conductivity to monitor peak elution during the SEC separation. If the salt were not removed, the conductivity signal of the impurities would be masked by the signal from the salt, and conductivity would only be useful for monitoring gross system changes.

10 The alkyl amines and/or alkyl ammonium compounds present in the described impurity apparently act as a counter ion to the phosphodiesters and/or associated to the polar portions of the triester groups of the antibacterial oligonucleotides. The impurity material can not be isolated from blank runs of 15 solutions, reagents, and stationary phases used during the described synthesis and purification procedures. Presently, the impurity has only been observed in oligonucleotides that have been AX purified.

Further characterization (by spectroscopic analysis) of 20 the stimulatory impurities isolated during the SEC step revealed that they are apparently produced by cleavage of absorption sites on the SAX stationary phase.

Although relatively crude oligonucleotide preparations were able to demonstrate significant inhibition in this assay 25 (after substantial removal of the contaminants that normally hinder the antibacterial effects of oligonucleotides), FDA requirements for parenteral therapeutics necessitate higher levels of purification for animal and human use.

30 5.6. Antigene Antibacterial Oligonucleotide Activity
Antibacterial oligonucleotides 96ss (SEQ ID NO. 79) and
73ss (SEQ ID NO. 124) (the ss denotes that oligonucleotide 73
is targeted to the sense strand) are homologous to the sense
strand of the targeted sequences. Oligonucleotides 96ss and
35 73ss are thought to exert antibacterial activity by acting as
antigene sequences that block gene expression by forming a
triple-stranded complex (i.e., triplex formation), or,

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possibly, by directly interacting with bacterial proteins. A time course of the antibacterial activity of oligonucleotides 73ss and 96ss is shown in Table 7.

5 5.7. The Use of Antibacterial Oligonucleotides Against Antibiotic Resistant Bacteria

The presently described antibacterial oligonucleotides are also capable of inhibiting the growth of a variety of bacteria that are known to be resistant to various traditional antibiotics. Tables 8(A-C and F) test the inhibitory activity of oligonucleotide 73 (NBT 73 - SEQ ID NO. 124) against clinical isolates of Escherichia coli that are known to be resistant to: streptomycin (8A); sulfonamide (8B); penicillin (8C); as well as multiple drug resistant Escherichia coli (8F). Oligonucleotide 114 (SEQ ID NO. 112) also inhibited the growth of Salmonella typhimurium ATCC accession No. 23564 (8D), Klebsiella pneumoniae ATCC accession No. 4352 (8E), and Staphylococcus aureus ATCC accession No. 29213 (8G).

Tables 9(A-G) test the inhibitory activity of oligonucleotide 114 (NBT 114 - SEQ ID NO. 112) against clinical isolates of Escherichia coli that are known to be resistant to: streptomycin (9A); sulfonamide (9B); penicillin (9C); as well as multiple drug resistant Escherichia coli (9F). Oligonucleotide 114 (SEQ ID NO. 112) also inhibited the growth of Salmonella typhimurium ATCC accession No. 23564 (9D), Klebsiella pneumoniae ATCC accession No. 4352 (9E), and

Staphylococcus aureus ATCC accession No. 29213 (9G).

Additional studies revealed that antibacterial oligonucleotides 114 (SEQ ID NO. 112), 5 (SEQ ID NO. 152), 39 (SEQ ID NO. 30), 43 (SEQ ID NO. 34), 3 (SEQ ID NO. 51), 78 (SEQ ID NO. 134), 12 (SEQ ID NO. 153), 14 (SEQ ID NO. 154), 23 (SEQ ID NO. 158), 24 (SEQ ID NO. 159), 22 (SEQ ID NO. 157), 17 (SEQ ID NO. 83), 20 (SEQ ID NO. 84), 15 (SEQ ID NO. 35 81), 16 (SEQ ID NO. 82), 19 (SEQ ID NO. 66), 28 (SEQ ID NO. 96), 63 (SEQ ID NO. 130), 10 (SEQ ID NO. 17), and 18 (SEQ ID NO. 73) significantly inhibited the growth of multiple drug

resistant Escherichia coli ATCC accession No. 35218 for over 400 minutes when present at a concentration of about 0.5-2.0 mg/ml as shown in Figures 6(a-t).

Additionally, antibacterial oligonucleotides 16 (SEQ ID 5 NO. 82), 18 (SEQ ID NO. 73), 1 (SEQ ID NO. 119), 5 (SEQ ID NO. 152), 17 (SEQ ID NO. 83), 21 (SEQ ID NO. 156), 132 (SEQ ID NO. 15), 11 (SEQ ID NO. 18), 89 (SEQ ID NO. 61), and 2 (SEQ ID NO. 50) all inhibited the growth of penicillin resistant clinical isolates of Staphylococcus aureus ATCC accession No. 13301 for over 400 minutes when present in the culture medium at a concentration of about 0.5-2.0 mg/ml (data are respectively provided in Figures 7(a-j)).

Oligonucleotide 14 (NBT 14 - SEQ ID NO. 154) was used to test whether the antibacterial oligonucleotides could also be 15 used to enhance a target bacteria's sensitivity to antibiotics to which the bacteria had previously proven resistant. Table 10 shows the results of a growth inhibition time course experiment where oligonucleotide 14 was tested for the ability to inhibit the growth of Escherichia coli Y1088 (known to be resistant to ampicillin) in the presence and absence of the indicated concentration of ampicillin (50 µg/ml, and 250 µg/ml). Table 10 indicates that oligonucleotide 14 is capable of significantly restoring ampicillin sensitivity of Escherichia coli Y1088.

5.8. <u>Animal Studies</u>

Preliminary assessments of the *in vivo* efficacy of the presently described antibacterial oligonucleotides (using a Lister & Saunders test) indicate that a higher percentage of animals treated with oligonucleotide survive exposure to near-lethal amounts of *Escherichia coli* ATCC accession No. 25922 (prepared and injected as described in Lister & Saunders, 1995). In particular, Figure 8 shows that mice treated with oligonucleotide 114 (SEQ ID NO. 112) *in vivo* 35 proved more resistant to challenge by a bacterial pathogen than control animals. The assay was conducted essentially as described in section 4.6, *supra*, and involved a total of 5 mg

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of oligonucleotide injected (I.P.) over a 2 day period (1 mg of oligonucleotide suspended in 0.5 ml of sterile saline was injected at 1, 5, 10, 24, and 34 hours post infection). Additionally, Figure 9 shows that mice treated with the 5 antibacterial oligonucleotide SOT 114.21 (phosphorothioate GGAACGCGC linked to 2'-methoxy riboCATTGGTATATC with an inverted 3' terminal deoxythymidine) had substantially enhanced survival after challenge with lethal quantities (approximately 10⁸ cfu in mucin and iron dextran injected i.p. 10 into CD1 mice) of Staph. Aureus. In Figure 9, treatment with Staph. was T=0 and 5 hours after infection. Oligonucleotide treatment was only administered on day 1.

Subsequent in vivo studies have shown that SOT 114.21 can increase the survival of Staph. Aureus challenged test 15 animals by about 81 percent, and increase the survival of E. coli infected test animals by about 95 percent (relative to animals treated with a placebo).

Similarly, when a representative antibacterial oligonucleotide was tested using the model of Hof et al.,

20 additional evidence of in vivo efficacy was obtained. In particular, Table 11 shows that mice treated with oligonucleotide 132 (SEQ ID NO. 15) in vivo had markedly reduced amounts of bacteremia 24 hours after initial exposure to Escherichia coli ATCC accession No. 25922 (prepared and injected as described in Hof et al., 1986). This assay was conducted essentially as described in section 4.6, and involved the injection of a total of 2 mg of oligonucleotide (1 mg injected at 6 and 10 hours post infection).

5.9. Standard MIC Assays

To eliminate the possibility that the observed antibacterial activity might be a function of the slightly 5 modified version of the MIC used to generate the above data, antibacterial assays were conducted using the standard MIC assay. Given that 44 percent of all nosocomial infections are caused by Staph. aureus, Streptococcus, or Pseudomonas, these bacteria were used as targets for standard MIC assays.

In brief, the standard MIC assay was conducted by using 10x13 mm tubes to which 40 μ l of Mueller Hinton Broth (purchased from BBL, obtained through VWR, 3745 Bayshore Blvd., Brisbane, CA 94005) was added. The oligonucleotides (including an oligo dT control) were supplied as lyophilized 15 pellets and dissolved in 200 μ l of sterile tissue culture water (Sigma), and 200 μ l aliquots of water or dissolved oligonucleotide were then added to the "control" or "oligo test" tubes.

Bacterial suspensions were prepared by suspending the 20 organisms in 1.0 ml of sterile-filtered saline (Sigma) at a concentration corresponding to an $O.D._{625}$ of 0.1-0.102. Ten μl of this suspension was then added to 990 μl of saline and 500 ul of this mixture was added to both the "control" and "oligo test" tubes (a concentration of approximately 1×10^5

- 25 bacteria per ml). Sterile saline was added (260 μl) to each of test tube to reach a total volume of 1 ml, the tubes were vortexed, O.D.₆₂₅'s were measured (time zero), and tubes were incubated at 35° C for 16-24 hours (without shaking). Tubes were vortexed in the morning, and the amount of bacterial 30 growth (if any) was measured by measuring O.D.₆₂₅ readings.
- Results from studies using the standard MIC assay are described in Figures 10 through 13.

The antibacterial oligonucleotides used in the following studies were constructed as follows (5' to 3'):

SOT T12, 12 thymidines (first six bases phosphorothicate deoxynucleotides, followed by six 2'-methoxy ribonucleotides and an inverted 3' terminal deoxythymidine linked by a 3'-3'

phosphodiester linkage); SOT-C12, 12 cytidines (first six bases phosphorothioate deoxynucleotides, followed by six 2'-methoxy ribonucleotides and an inverted 3' terminal deoxythymidine); SOT 89.6 (phosphorothioate deoxyCAT linked

- 5 to 2'-methoxy riboGTC with an inverted 3' terminal deoxythymidine); SOT 89.9 (phosphorothioate deoxyCATGT linked to 2'-methoxy riboCATT with an inverted 3' terminal deoxythymidine); SOT 89.12 (phosphorothioate deoxyCATGTC linked to 2'-methoxy riboATTCTC with an inverted 3' terminal
- 10 deoxythymidine); SOC 89.12 (phosphorothioate deoxyCATGTC linked to 2'-methoxy riboATTCTC with a 3' terminal cholesteryl group); SOB 89.12 (phosphorothioate deoxyCATGTC linked to 2'-methoxy riboATTCTC with a 3' terminal biotin group); MMT 89.12 (89.12 with all methoxyribonucleotides
- 15 linked to an inverted 3' terminal deoxythymidine); MPT 89.12 (the 89.12 sequence, CATGTCATTCTC, with all p-ethoxy, 2'-methoxy RNA linked to an inverted 3' terminal deoxythymidine); SOPT 89.12 (phosphorothioate deoxyCATGTC linked to 2'-methoxy riboATTC followed by p-ethoxy, 2'-
- 20 methoxy riboTC linked to an inverted 3' terminal
 deoxythymidine); POT 89.12 (89.12 with all p-ethoxy DNA
 linked to an inverted 3' terminal deoxythymidine); DSM 89.18
 (phosphorothioate deoxyCATGTCAT linked to phosphorothio
 (i.e., sulphur), 2'-methoxyriboTCTCCTTAAG linked to a 3'-
- 25 terminal deoxythymidine); SSM 89.18 (sulphur, 2'-methoxy
 riboCATGTCATTCTCCTTAAG linked to a 3'-terminal
 deoxythymidine); NBT 89.15 (phosphorothioate deoxy
 CATGTCATTCTCCTT linked to an inverted 3' terminal
 deoxythymidine); NBPT 89.12 (phosphorothioate deoxyCATGTC,
- 30 linked to 2'-methoxy riboATTC, followed by p-ethoxy, 2'methoxy riboTC linked to an inverted 3' terminal
 deoxythymidine); MMPT 89.12 (2'-methoxy riboCATGTCATTC linked
 to p-ethoxy, 2'-methoxy riboTC, linked to an inverted 3'
 terminal deoxythymidine); SST 89.12 (phosphorothioate
- 35 deoxyCATGT linked to sulphur, 2'-methoxy riboCATTCTC linked to an inverted 3' terminal deoxythymidine); SOT 1.15 (phosphorothioate deoxyTGTGTA, linked to 2'-

methoxyriboGCCCATAGT, linked to an inverted 3' terminal deoxythymidine); SOT 5 (phosphorothioate deoxyTTGAC linked to 2'-methoxy riboATATCGGTCACTC linked to an inverted 3' terminal deoxythymidine); SOT 143.15 (phosphorothioate

- 5 deoxyCTCATG linked to 2'-methoxyriboATTAACACC linked to an inverted 3' terminal deoxythymidine); SOM-89 (a sulphur, 2'-methoxyriboC, linked to phosphorothioate deoxyGCCA, linked to 2'-methoxyriboTGTCATTCTCCT, linked to sulphur, 2'-methoxyriboTAA, linked to a 3' terminal deoxyguanidine); SOM
- 10 72.1 (a 5' sulphur, 2'-methoxyriboA, linked to
 phosphorothioate deoxyCTGA, linked to 2' methoxyriboTGACTTCATGAT, linked to sulphur, 2' methoxyriboGCG, linked to a 3' terminal deoxycytosine); SOT
 89.21 (phosphorothioate deoxyCGCCATGT linked to 2'-
- 15 methoxyriboCATTCTCCTTAAG linked to an inverted 3' terminal deoxythymidine), SOM 114 (phosphorothioate deoxyGGAACGCG, linked to 2'-methoxyriboCCATTGGTA, linked to sulphur, 2'methoxyriboTAT, linked to a 3' terminal deoxycytidine), MMT 89.12 (2'-methoxyriboCATGTCATTCTC linked to an inverted 3'
- 20 terminal deoxythymidine); 132 (SEQ ID NO. 15), SOM 1.1 (sulphur, 2'-methoxyriboA, linked to phosphorothioate deoxyGCAA, linked to 2'-methoxyriboCTGTGTAGCCCA, linked to sulphur, 2'-methoxyriboTAG, linked to a 3' terminal deoxythymidine, SOM 72.1, or SOM 5.1 (sulphur, 2'-methoxyT,
- 25 linked to phosphorothioate deoxyACTT, linked to 2'methoxyriboGACATATCGGTC, linked to sulphur, 2'methoxyriboACT, linked to a 3' terminal deoxycytidine), and
 mixtures of SOT(5.15, 78.15, 89.15, and 114.15) or SOT(89.18,
 114.15 (phosphorothioate deoxyCGCCAT linked to 2'-
- 30 methoxyriboTGGTATATC linked to an inverted 3' terminal deoxythymidine), and 78.15 (phosphorothioate deoxyCATTGT linked to 2'-methoxyriboTTGTACTCC linked to an inverted 3' terminal deoxythymidine).

Figures 10a and 10b show the results of standard

35 overnight MIC assays using the indicated oligonucleotides to test for antibacterial activity against Staph. aureus.

Virtually all of the oligonucleotides tested (SOT-T12, SOT-

C12, SOT 89.(6, 9, and 12), SOC 89.12, SST 89.12, SOT 1.15, SOT 5.15 (phosphorothioate deoxyACATAT linked to 2'-methoxyriboCGGTCACTC linked to an inverted 3' terminal deoxythymidine), and SOT 143.15) significantly inhibited the growth of Staph. aureus (with the exception of the oligo dT string) relative to the control samples.

Figures 11a and 11b show the antibacterial activity of oligonucleotides DSM 89.18, SOT 78.15 (phosphorothioate deoxyCATTGT linked to 2'-methoxyriboTTGTACTCC linked to an 10 inverted 3' terminal deoxythymidine), SOM 114.15, SOT 89.18 (phosphorthioate deoxyCATGTCAT linked to a 2'-methoxyriboTCTCCTTAAG, linked to an inverted 3' deoxythymidine), SOT 89.21, NBT 89.15, NBT 89.12-1 (phosphorothioate deoxyCATGTCATTCTC linked to a 3' terminal inverted phosphorothioate deoxythymidine), NMPT 89.12-2

- 15 inverted phosphorothioate deoxythymidine), NMPT 89.12-2 (phosphorothioate deoxyCATGTCATTC linked to 2'-methyl, pethoxy TC, linked to an inverted 3' terminal deoxythymidine); MPT 89.12-4 (CATGTCATTCTC, with all pethoxy, 2'-methoxy RNA linked to an inverted 3' terminal deoxythymidine); MMPT
- 20 89.12-5 (2'-methoxy riboCATGTCATTC linked to p-ethoxy, 2'methoxy riboTC, linked to an inverted 3' terminal
 deoxythymidine); SOT 89.12-6 (phosphorothioate deoxyCATGTC
 linked to 2'-methoxy riboATTCTC with an inverted 3' terminal
 deoxythymidine); SOPT 89.12-7 (phorphorothioate deoxyCATGTC
- 25 linked to 2'-methoxy riboATTC followed by p-ethoxy, 2'-methoxy riboTC linked to an inverted 3' terminal deoxythymidine) when measured in standard overnight MIC assays against Serratia liquefaciens. As is readily apparent, all of the test oligonucleotides displayed
- 30 significant antibacterial activity relative to controls.

 Interestingly, the oligonucleotides used in Figures 10-

11 retained antibacterial activity when used in Figures 10-11 retained antibacterial activity when used in standard overnight MIC assays over the three day time course. These data indicate that the tested antibacterial oligonucleotides 35 are bactericidal for the test microorganisms.

Figure 12 shows the level of growth inhibition obtained when the oligonucleotides SOC 89.12, SOB 89.12, MMT 89.12,

MPT 89.12, SOPT 89.12, POT 89.12, DSM 89.18, SSM 89.18, NBT 89.15, NBPT 89.12, MMPT 89.12, SOT 89.12, and SOM-89Filwere tested in a standard MIC assay against Staph. aureus. All of the tested oligonucleotides proved effective at inhibiting 5 the growth of Staph. aureus.

Figure 13 shows that several different length variants meths@Tr89o21 [60k±2,t150](phosphorothioate deoxyCATGTC linked to 2'-methoxy riboATTCTCCTT linked to an inverted 3' terminal deoxythymidine), and 18mers) were able to inhibit the growth 10 of Staph. aureus when they were tested in a standard MIC assay against Staph. aureus.

Figures 14(a and b) compare the antibacterial activities of the conventional antibiotic ampicillin and SOT 114.21 against isolates of Staph. aureus strains 13301 and 29213.

15 Figure 15 shows that oligonucleotide MMT 114.15 (2'imethoxyriboCGCCATTGGTATATC'linked to an inverted 3' terminal
deoxythymidine) proved capable of inhibiting the growth of P.
aeroginosa strain 10145, an opportunistic Gram negative
pathogen that has proved resistant to many conventional
200 antibiotics, in a standard MIC assay.

Figure 16 shows that oligonucleotide SOT 114.21 proved capable of inhibiting the growth of the pathogen *Strep*.

pyogenes strain 14289 in a standard MIC assay.

25 EQUIVALENTS

The foregoing specification is considered to be sufficient to enable one skilled in the art to broadly practice the invention. Indeed, various modifications of the above-described methods for carrying out the invention which 30 are obvious to those skilled in the field of microbiology, biochemistry, organic chemistry, medicine or related fields are intended to be within the scope of the following claims. All patents, patent applications, and publications cited are incorporated herein by reference.

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TABLE 1 Table 1 Antimicrobial Susceptibility According to NCCLS Standards Gene/ NBT Drug R Drug R Gram Neg Gram Pos SEQ Sequence Operon Target Category of Target: Energy Metabolism 97% INH 100% INH 1 PAG GGT CAT GTC TGC GGG AAA TAA TAC NBT 28 hema NBT 32 57% INH 2 CCG TTA TTG TTG TGT TTG CGT GTT TAC NBT 36 aroA 3 CAG GGA TTC CAT GAA ACT CAA CTC TCA chaC NBT 47 ACA CTT CCG CCA CTG CAT ACT TCC CTG 4 chaB NBT 48 5 TCG TTT TAT ACG GCA TCG TTG ACT CCT chaA NBT 49 6 GAC ATT ATG GTT ATC CCT TTG CAG ATG ATP NBT 57 S64 INH TTC ACT CCT GCT CCC TTC GAG GTA TGC operon NBT 61 hemD 8 GCG GGT GAC AAG GAT ACT CAT GCC GGG NBT 62 hemX 9 CAT TAT GGC TTC CTG TTA TGA GAG TTA son NBT 67 GTT GTG AAG CCA TGT ACA CCT TTC CAG operon NBT 84 78% INH 26% INH crp 11 GTT TGC CAA GCA CCA TGC GCG GTT TAC ATPase NBT 88 72% INH 12 CGT CAT ATT TTC TGA AGC CAT GAT GCC NBT 104 суа GGT ACA AGA CGT ATC GCC TGA TTT GCT pckA NBT 126 14 CAT TTC TCA GCT CCT TAG CCA ATA TGT NBT 132 89% INH 100% INH fadD 15 AGC CAA ACC TTC TTC AAT TCT TCA CCT Category of Target: DNA Replication NBT 9 NBT 10 100% INH 33% INH ATVP 16 17 AAG GTC GCT CAT CTA ACC GCT ATC CCT AGG TAA TTC AGC CAT CAA GAG TTC CTC NBT 11 gyrB 96% INH 100% INH 18 AAT GCA GTC ACC ATC GCT TTC TGT TAC lig NBT 26 19 GCA TCA GCC TGT CGT ATT CAG CGT CGG dnaG NBT 30 20 CGG CTC GTT TTC ACG TAC TTT AAT TAC ssb NBT 37 21 TCT GCT GGC CAT AAT TGA GTC TCC TGA groESL NBT 66 63% INH 22 ATA ACT CTC CTT TGA GAA AGT CCG TAT dna A NBT 79 65% INH 23 AAG CGA AAG TGA CAC GGC GGA CTC CAC operon dnaT 24 GGT CAT CAA GAT CAT TCG GGA ACC ATG NBT 81 operon parc NBT 95 25 TCG CTC ATT AAT TCT GAT TCC TCA ACT holD NBT 109 26 TAA CTG CCA GTC TCG TCG GGA TGT CAT dnaQ NBT 124 27 CGT GTA ATT GCA GTG CTC ATA GCG GTC dnaE NBT 130 28 TGT ACG AAA CGT GGT TCA GAC ATC TTC dnaJ NBT 133 29 CTC GTA ATA ATC TTG CTT AGC CAT CTT Category of Target: Cell Division Control NBT 39 100% INH 34% INH minB 30 TGA CAT CCT GGC CTT ACT CAA TTA GCT minD NBT 40 CAA CAA TAA TGC GTG CCA TAG AAA TTC minE NBT 41 12 GAG TAA TGC CAT AAC TTA TCC TCC GAA 33 AAC GCA TCA ACC TAA CTC CTT CGC CAG ftsW NBT 42 ftsN NBT 43 100% INH 34 TAT TTA TTC GTT CGT CAG CCC GCC ATG

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			And Susceptibil to MC	imicrobial lity Accordin CLS Standard	ig is	
	ftsH	NBT 44			35	GAA CGC TTC TTA CCT GTC ATT TAA CT
5	ftsJ	NBT 45			36	GTA TTA GGT TTT TCG CCA TGT CAC TCA
5	ftsQ	NBT 52			37	TTC AGA GCA GCC TGC GAC ATA TTA GAC
	ftsA	NBT 53			38	CGT CGC CTT GAT CAT TGT TGT TCT GCC
	ftsZ	NBT 54			39	ATT GGT TCA AAC ATA GTT TCT CTC CGA
	ратВ	NBT 55	<u> </u>	. _	40	TGG ATG TIT CAT GGC CTT CTC CTT
	fts YEX operon	NBT 65	60% INH	21% INH	41	CTA CAC TCC TCG CTG TTG CTT CAT GGC
10	рьрв	NBT 80	90% INH	42% INH	42	TGC TTT CAT GCG TCG CGT TTA TCC TTA
	rodA	NBT 83	 	<u> </u>	43	TTA TCC GTC ATG ATT AAT GGT CCT CCG
	tig	NBT 119	<u> </u>	<u> </u>	44	GCA TCT TGT TAC CTC AAA AAA TCA CAG
	Categor	y of Tax	rget: Re	gulatory F	rote	ina
	lon	NBT 27	97% INH	100% INH	45	AGG ATT CAT AGA GCT CTC TAG TTT
	rel B	NBT 56	ļ	ļ	46	TTA ACA TCT TTT GCT GCT GCT TCA TAG
15	crp	NBT 84	78% INH	26% INH	47	GTT TGC CAA GCA CCA TGC GCG GTT TAC
	lexA	NBT 131	<u> </u>		48	GCC TGG CCG TTA ACG CTT TCA TTC CGC
	Categor	of Tar	rget: Ce	11 Wall Bi	OSYM	thesis
		NBT 1 NBT 2 NBT 3 NBT 7 NBT 8	97% INH	100% INH	50 51 52	AAC GAT AGC AAC TGT GTA GCC CAT AGT AGG AGT TCC ACC GGC GCC TGT AGC ACC GGA GCC GAT CAA CAA CTG CGA TGG TGG TGA TTG TAA ACC GCG CCA GCC GAC GAC GAA ACC TAC TTT
20	11 1	NBT 33 NBT 34 NBT 35	70% INH	26% INH	54 55	CGT CTA ACA CAA AGT GCA TAC ATT ACC CAC TTC ATG TTG CGC TGA TTT ACC ACC CCA TCT TAA AAA CCT ATC CCG TCT AAC
	murG	NBT 50		<u> </u>	57	CTT GAC CAC TCA TCG TGA ACC TCG TAC
	murc	NBT 51	<u> </u>		58	GTT GTG TAT TCA TTC TTT ACG CCA TTA
	lysA	NBT 6	<u> </u>		59	CAG TGA ATG TGG CAT AAC AAA CTC CAG
	murD	NBT 139	<u></u>	<u> </u>	60	CGT ACC TTC AGC GTT GCC AGA CCA ATC
25	Category	of Tar	get: Sug	ar Ketabo	lism	
	zwf	NBT 89	69% INH	100% INH	61	CGT TAC CGC CAT GTC ATT CTC CTT AAG
	sdhB	NBT 112			62	CTG TCT CCG CAT TAG TAA GTA CGA ATC
	Category	of Tar	get: Vir	ulence, P	111,	Plagalla
	pap operon	NBT 72	31% INH		63	TGA CCG ACT GAT GAC TTC ATG ATG CGC
	rim J h	NBT 103	76% INH		64	CAT TOT ATA COT ACT COT TOC CGT AAC
30	Category	of Tar	get: Pat	ty Acid Ne	tabo	lism
	fadD 1	NBT 132	89% INH	1001 INH	65	AGC CAA ACC TTC TTC AAT TCT TCA CCT
	Category	of Tar	get: mRN	A Synthesi	s/St	ability
	rpoN N	/BT 19	100% INH	23 * INH	66	TAG GAT GTT CTA ACC TTT TCA ATC AGC
	alpha h operon	IBT 29	98 % INH		67	IAC GGG CCA CTA TGC ACT CCT ACT ATT
35	MMS N operon	BT 30			68	CGG CTC GTT ITC ACG TAC TTT AAT TAC
	rho N	BT 125			69 K	GAT TCA TAG TGG TGT GAG TTC TTA AAC
	rnpB N	BT 121			70 C	GAA GAG GAC GAC GAA GCG GCG ACT

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	ļ							Tabl	e 1			·					
			5	iuscep	Anci Cibili CO NC	aicro ty Ac LS St	biel cordin andero	7									
	ams	NBT	134	<u>L</u>				71	TCA	TCC	TA	CT	' AC	CAT	TAT	TCT	TAC
5	Catego	ry of	Tax	get:	tRi	A Sy	nthe	1.									
J	trm D	NBT	16	100	INH	100	* INH	72	GCI	AAA	CGA	ATA	GT	r acc	ATA	ACA	TCC
	met Y	1 10	18 8.12 8.15 8.18	100	inh	100	* INH	174 175	LCY LCY	TCT	CIG	CTA	ATT	ATT		CTC	TAA
	val U	NBT NBT	91			\vdash		74 75	GTG		TCC	CAG	CTC	AGC	TAA		
0	tRNA operon	NBT NBT	93					76 77	CGC	TCT	ATC	CAG	CTG	AGC	TAC	GGG	
	infA operon	NBT						78	1					TAC			ATT
	Categor	yot	Tar	get:	ran	A Sy	nthes	is	1								
	rrn8 operon	NBT	96	80%				1	GCC	GCC	AGC	GTT	CAA	TCT	GAG	TGA	
	Categor	y of	Tar	get:	Rib	osom.	al Pr	oteli	Sy	nthe	eis	-					
5	str operon	NBT		971				1				CAT	TAA	ATA	GCT	CCT	GGA
	slO operon	NBT	15	1001	HNI	40%	INH	81	GCG	GAT	ACG	GAT	TCT	TTG	GTT	cre	CAT
	trmD operon	NBT	16	100%	HNI	100%	INH	82	GCT	AAA	CGA	ATA	GTT	ACC	ATA	ACA	TCC
	spc operon	NBT		100%	INH	981	INH	83	GTT	CAG	CAT	AGT	CTG	TTC	TTG	GAT	CAT
, 	S15 Operon	NBT		991			INH							ACT			
	S12 operon	NBT	21	821	INH	100%	INH	85	TTG	TAG	GCA	זכד	ACA	TTC	TCC	TGT	CII
	alpha operon	NBT	-	981	INH			86	TAC	GGG	CCA	CTA	TGC	ACT	CCT	ACT	ATT
	MMS operon	NBT .						87	CGG	стс	GTT	TTC	ACG	TAC	111	AAT	TAC
'	tsf.	NBT		671		461	INH	88	AAA	CAG	TTG	CCA	TGA	ATT	TTT	CCT	CTA
	rim J	NBT		781	HNI			I						CCT			
	rim I	NBT :												TCG TAA			
	rnpA	NBT :												CAG			
	rpmH .	NBT I							GCG	TTT	CAT	GGC	GAT	TTC	TAC	CTA	AAC
	Str operon	NBT 1	Ť	97 %]	$\neg \neg$	•in	Synth		AAC	TGT	TGC	CAT	TAA	ATA	GCT	CCT	GGA
		NBT 1	18 2	100	INH	1001	INH	95	TAA '	TCA	TCT	CIG	CTA	ATT	17G	crc	TAA
		NBT 2	8 9	7¥ I	NH :	1001	INH	96	LAG (GGT	CAT	GTC	TGC	GGG	AAA '	TAA '	TAC
	infC	NBT 3	1					- 1						TTA			
į	tsf	NBT 3	8 1	001	INH 4	61 1	NH	- 1						TTA			
	prf	NBT 9	0		\Box									AAT '			 ()
	infA operon	NBT 1	10					100									
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	<u> </u>				Tabl	le 1						
			An Susceptibi to M	timicrobial lity Accordi TCLS Standar	ng ds							
	aat	NBT 1			- I	TAT TO	A ATG	GCG A	GA AA	G CA	S AA	CAC
_	Catego	ry of T	arget: Pl	ospholipi	d Syn	thesis						
5	adk	NBT 25	96% INH	44% INH	102	CCG AA	G CAG	AAT A	AT AC	G CAT	TAC	: GA
	psd	NBT 10	5 831 INH	SOT INH	103	AAA TG	A ATT	TAA C	AA GG	T AGO	· cro	CAG
	pss	NBT 10	6		104	CAG TG	CATT	TCT T	CT CT	3 770	ATT	GAA
	Catego	ry of T	rget: Pe	riplasmic	/Secr	etory P	rotei	3.6				
	envA	NBT 46	591 INH	25% INH	105	TTT GT	T TGA	TCA T	CG TA	TAT	CTC	GCC
	tolA	NBT 85			106	CGG TT	CCT	TTG A	CA CT	TCC	GTI	TCC
10	tolB	NBT 86			107	CCT GC	T TCA	TCA T	AT CT	CCT	ATA	CTG
	secA	NBT 11	8	_i	108	CTT TA	J TTA	ACA A	IT TG	TTA	GCA	TAA
	Catego	ry of Ta	rget: Tr	ansport Pr	rotei	n.e					,	
	biotin	NBT 58	84% INH		109	GCG AC	ATG	TCC AC	C GT	GGC	GGT	GAG
	operon	NBT 59 NBT 60				GTT AAT						
	fhuA	NBT 11	100% IN	18% INH	112	GGA ACC	ccc	CAT TO	G TA1	ATC	TCT	GAT
15	fhuC	NBT 11	5		113	TCC TGC	ATA	ACA GO	C AAC	TTG	TGA	TTA
•	jhuD	NBT 11	5	<u> </u>	114	TAA GAC	GTA	AGC CC	C TC	TCA	ATA	AAC
	fhuB	NBT 11	,		115	CTG CG	GAA	GTT_CA	T CCA	GGT	GAG	CGC
	Categor	y of Ta	rget: Am	ino Acid B	iosyr	thesis						
	aroC	NBT 32	97% INH	<u> </u>	116	CCG TTA	TTG	TTG TG	TTTG	CGT	GTT	TAC
	aroA	NBT 36			117	CAG GGA	TTC	CAT GA	A ACT	CAA	CTC	TCA
20	nir operon	NBT 71	93 % INH	42% INH	118	ATA ATT	GCG	AGT CT	G ACT	TTG	CTC	ATT
	asd	NBT 1 NBT 2 NBT 3 NBT 7 NBT 8	97% INH	100% INH	120 121 122	AAC GAT ACG AGT GGA GCC CAA CAA ACC GCG	GAC CTG	ACC GG CAT AC CGA TG	C GCC C GCG G TGG	TCT CCA TCA	AGC GCC TTG	ACC GAT TAA
	Categor	y of Ta	get: Lip	opolysacc								
25	rfaY	NBT 73	1001 INH	J6% INH	124	GTC TIT	GAT (CTT GC	T CTT	CTG	AAT	CAT
2.5	rfaZ	NBT 74				TAT CTA						
	rfaL	NBT 75				TTC CTA					CAT	
	rfaK	NBT 76			127	TAA TGA	TGA 1	TAA CT	TTTC	CAA	AAC	TGC
	l ps operon	NBT 77	76% INH		128	CCA TGA	TAT (GC AT	C TTT	ATG	ACC	AGG
	Categor	y of Tax	get: Pur	ine/Pyrim	idine	Biosyn	thesi	,				
30	adk	NBT 25	96* INH		129	CCG AAG	CAG 2	AA TAJ	r acg	CAT	TAC	GAA
	deoC operon	NBT 63	1001 INH	51% INH		GCT TTC						_
	pyrE operon	NBT 64			131	ITC ATC	ATA A	CG GG	CAC	GAT	CTC	GTC
	prs	NBT 111	87% INH		132	CAT ATC	AGG C	AC CAC	AAG	AAC	CTC	AGG
		NBT 128				TTC GCT						
35		of Tax	get: Out	er Hembrar								
	i	NBT 78	100\$ INH			STA GTT	стс т	TG CAT	TGT	TTG	TAC	TCC
		NBT 87			135	GC TAG	ATG A	TG TG1	тст	CAG	TTT	CAT

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	<u> </u>					-		Tab	le :	L							
				Susci	epcibi.	Lity A	robial Accordi	_g ds			_	-					
	отрх	NE	T 97					13	6 CA	T AA	C CA	C CT	C AA	A TG	T G	T TC	A AA1
5	ompF	NE	T 98						- 1			-					T ATT
3	опрС	NB	T 99														T GT1
	Оперн	NB	T 10	•													C ATT
	Omp P	NB	T 10	1		1											T GAC
	OmpA	NB	T 10	2		1										T CA	
	tsx	NB	T 12						- 1								A TCC
3.0	1pp	NB	T 12	7 911	INH	981	INH	1									r CAT
10	envM	NB	T 129	ч_	_	\perp											TGA
	envC	NB	T 13	<u> </u>		↓_											CAA
	envD	NB.	T 136	4		1_			- 1								TTC
	envR	NB:	T 136	Ц				147	111	GCC	ATG	ATI	` אאז	TAT	TC	(GG2	AAT
	Categor	y o	f Ta	rget	. NT	trate	Red	cta	e .								
15	nar operon	NB7	T 68	70%	INH	_		148	ATT	TAC	TCA	TCG	GTT	TTC	TCC	: TGT	GGC
	nar XL operon	NBT	F 69	\perp				149	AAG	CAT	GTA	AAC	CIC	TTC	C::1	CAG	GCT
	nar ZYWZ operon	NBT	70					150	GAT	CCA	AAA	GTT	TAC	TCA	TAG	CAT	GAC
	nir operon	NBT	71	80%	INH	42%	INH	151	ATA	ATT	GCG	AGT	CTG	ACT	175	CIC	ATT
20	Categor	y 0:	Tax	get:	Dry	g Re	mista	DC 6									\neg
•	sulA	NBT	· 5	100	INH	1001	INH	152	TGG	CTT	TAC	TTG	ACA	TAT	ന്നു	TCA	СТС
	str operon	NBT	12					ł								ССТ	
	bla	NBT	14	994	INH	981	INH	154	ACA	CGG	AAA	TGT	TGA	ATA	CIC	ATA	CTC
	spc operon	NBT	17	1001	INH	984	INH	1	1							GAT	
25	S12 operon	NBT	21	821	INH	1001	INH	156	TTG	TAG	GCX	TCT	ACA	TTC	TCC	TGT	GTT
	tet resista nce	NBT	22	100	INH	901	INH	157	ATT	GTT	AGA	TTT	CAT	ACA	CGG	TGC	CTG
	kan resista nce	NBT	23	98%	INH	104	INH	158	CAT	CTT	GIT	CAA	TCA	TGC	Gλλ	ACG	ATC
30	ermC	NBT	24					159	ACT	GTG	TIT	TAT	ATT	TTT	כייכ	GTT	CAT
30	рърв	NBT	80	901	INH	421	INH									TCC	
	PbpA	VBT	82												_	GCT	
	Category	of	Targ	et:	Vita	min											
-	biotin operon	VBT	59	841	INH			101 F	ATC	GGG	ctt :	~~	ממח	B B T	3	GGT TTG	[i
35		VBT	\rightarrow	1001	ниі	1001	INH	164	311	AAT	TCG	GTG '	TAG A	ACT	TOT	AAA	CCT
_	Category	~*			<u> </u>	<u> </u>											$-\parallel$
1.1			\neg			9118		,,,									
. Ш	<u></u>	D1	144	.001	TMH			166	CI	LAT	CAA A	ACA Z	ATG				

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Table 1 Antimicrobial
Susceptibility According
to NCCLS Standards NBT 140 100% INH 167 ATA TAT ATA TAT ATA TAT NBT 141 100% INH (AC), 168 ACA CAC ACA CAC ACA CAC (TC), NBT 142 100% INH 169 TCT CTC TCT CTC TCT CTC (T), NBT 13 100% INH 170 TTT TTT TTT TTT TTT (C), NBT 143 100% INH 171 | CCC CCC CCC CCC CCC NBT 113 sucA 172 CAA AGC GCT GTT CTG CAT CGT GAT CCC (RS) NBT 4 173 GAT ATC CGC ATG GTT CAA CAG ATG ACA

10

5

15

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25

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	Table 2A.	Energy Mecabolist	<u> - Oligonucleotid</u>	e #28						
	Escherichia coli 35218 Multiple Drug Resistance									
	Time T=0	Control 0	28	*Inhib						
5	0	0	0							
	60	0.002	0.001	50%						
	120	0.001	0.001	01						
	180	0.003	0	100%						
- 1	240	0.008		100%						
10	285	0.015	0	1001						
_	320	0.026	0	100%						
	350	0.04	0	100%						
	380	0.058	0.001	984						
- !	410	0.076	0.002	97%						
ļ	430	0.091	0.004	961						
15 [450	0.105	0.004	96%						

DNA Replication - Oligonucleotide #10 20 Escherichia coli 35218 Multiple Drug Resistance Time T=0 Control 10 *Inhib 0.003 0 0 0 60 0.001 0.001 08 25 120 0 0 170 0.003 0 1001 230 0.008 0 100% 275 0.017 0 100% 305 0.025 0 1001 340 0.046 0 100% 30 365 0.058 0 100% 385 0.075 -0.002 103% 400 0.082 -0.002 1024 415 0.094 -0.002 102% 425 0.105 0.001 991 35

0.001

0.002

0.008

991

981

93**%**

Table 2C. Cell Division Control - Oligonucleotide #43 Escherichia coli 35218 Multiple Drug Resistance Time T=0 Control *Inhib 0.005 5 0 0 0 105 -0.001 0.002 150% 175 0.003 -0.004 233% 220 0.004 175% -0.003 270 0.007 -0.003 1431 300 0.012 -0.003 1251 10 330 0.022 -0.003 1144 360 0.032 -0.002 106% 395 0.052 -0.001 102% 425 0.065 0 1001

0.081

0.09

0.108

445

465

490

	Table 20).	Regulatory Prote	ins - Oligo	nucleoti	de #27				
20	Escherichia coli 35218 Multiple Drug Resistance									
	Time T=0		Control 0	27	0.002	*Inhib				
		0	0		0					
		60	0.002		0.001	50%				
25	,	20	0.001		0.001	0%				
		80	, 0.003		0	100%				
	2	40	0.008		0	100%				
	2	85	0.015		0	100%				
	3	20	0.026		0	1001				
	3	50	0.04		-0.001	103%				
30	3	80	0.058		0.001	98%				
1	4	10	0.076		0.002	971				
	4	30	0.091		0.003	97%				
l	4	50	0.105		0.003	97%				

Table 2E. Cell Wall Biosynthesis - Oligonucleotide #2 Escherichia coli 35218 Multiple Drug Resistance Time T=0 Control * Inhib 0.002 5 0 0 0 105 0.002 -0.001 1504 175 0.003 -0.002 167% 220 0.004 -0.001 125% 270 0.007 -0.001 114* 0.012 300 -0.001 108% 10 330 0.022 -0.001 105% 360 0.032 0 100% 395 0.052 0 100% 425 0.065 0 100% 445 0.081 0.002 981 15 465 0.09 0.003 97% 490 0.108 0.008 93 **t**

	Table 2	F. Sugar Metabo	olism - Oligonucleoti	de #89						
20	Staphylococcus aureus 13301									
	Time T=0	Control 0	89 0.003	* Inhib						
		0	٥							
	90	0.002	-0.002	2001						
	150	0.004	-0.002	1501						
25	210	0.008	-0.002	125%						
	255	0.015	-0.002	1134						
	285	0.026	-0.001	104%						
	315	0.039	-0.001	1034						
	345	0.052	-0.001	1021						
30	375	0.073	-0.002	1034						
	395	0.08	-0.001	101%						
- 1	415	0.089	-0.002	102%						
Į	435	0.103	-0.002	1021						

	Table 2G. Vi	rulence, Pili,	Flagella - Oligonucl	eotide #103
		Escherichi Multiple Dr	a coli 35218 ug Resistance	
	Time T=0	Control 0	103	* Inhib
5		0	0	
	60	0.001	-0.001	200%
	120	0.002	-0.002	2001
	180	0.006	-0.001	1171
	215	0.012	-0.001	1081
	250	0.02		100%
10	285	0.031	0.001	974
	. 325	0.072	0.009	881
	355	0.085	0.015	821
	375	0.096	0.021	781
[395	0.108	0.026	76%

Table	2H.	Fatty	Acid	Metabolism	-	Oligonucleotide #132
						TOTAL WIND

			a coli 35218 ug Resistance	
20	Time T=0	Control 0	0.003	* Inhib
	0	<u> </u>	0	
	60	0.001	-0.003	400%
	120	0.004	-0.002	1501
	165	0.007	-0.003	1434
	205	0.018	-0.002	1111
25	235	0.028	-0.002	1071
	265	0.039	-0.001	1034
	295	0.063	0.003	95%
- 1	315	0.078	0.004	951
	335	0.093	0.009	90%
30	355	0.107	0.013	88%

	Table 2I. mR	NA Synthesis/St	ability - Oligonucle	otide #19
•		Escherichi Multiple Dr	a coli 35218 ug Resistance	
	Time T=0	Control -0.001	0.005	* Inhib
5		0	0	
_	60_	0.001	-0.001	2001
	150	0.002	-0.001	150t
	195	0.005	-0.001	1201
	245	0.013	-0.002	115%
	275	0.019	-0.001	1051
10	320	0.04	a	1001
	350	0.054	-0.002	1044
- 1	365	0.066	0	100%
	385	0.079	-0.002	1031
l	415	0.095	0.003	971
	430	0.105	0.001	905

	Table 7	J. tRNA Synthe	sis - Oligonucleotide	#16							
	Escherichia coli 35218 Multiple Drug Resistance										
20	Time T=0	Control 0	0.003	\ Inhib							
	0	0	0								
	60	0.001	-0.002	3001							
	120	0	-0.002								
	170	0.003	-0.002	1671							
25	230	0.008	-0.002	1251							
	275	0.017	-0.004	1241							
	305	0.025	-0.004	116%							
	340	0.046	-0.004	1091							
-	365	0.058	-0.004	1071							
30	385	0.075	-0.004	105%							
30	400	0.082	-0.004	105%							
	415	0.094	-0.004	1041							
	425	0.105	-0.002	1021							

WO 98/03533 PCT/US97/12961

	Table 2	K. rRNA Synthe:	sis - Oligonucleotide	#96				
	Escherichia coli 35218 Multiple Drug Resistance							
	Time T=0	Control 0	96 0.005	1 Inhib				
5	0	0	0					
	60	0.002	-0.002	200%				
	120	0.004	-0.005	2251				
	165	0.005	-0.004	180%				
	210	0.011	-0.003	127%				
	250	0.018	-0.002	1111				
10	275	0.025	-0.001	104%				
	305	0.037	0.003	921				
İ	340	0.056	0.013	77%				
	360	0.069	0.02	711				
	380	0.08	0.028	65%				
15	400	0.096	0.042	561				
[420	0.108	0.053	514				

					TEOLIGE 42.		
20	Escherichia coli 15218 Multiple Drug Resistance						
	Time T=0	Control 0	21	0.002	t Inhib		
	0	0		0			
	60	0.001		-0.003	400%		
	120	120 0.004		-0.002	1501		
25	165	0.007		-0.004	1574		
j	205	0.018		-0.001	1061		
ļ	235	0.028		-0.001	104%		
	265	0.039		0.001	971		
i	295	0.063		0.007	891		
30	315	0.078		0.01	87%		
- 1	335	0.093		0.018	814		
(355	0.107		0.025	771		

Table 2M. Protein Synthesis - Oligonucleotide #18 Escherichia coli 35218 Multiple Drug Resistance Control 0.001 Time T=0 * Inhib 0.017 0 0 0 5 .60 0.001 -0.004 500% 120 0.002 -0.004 300% 165 0.005 -0.009 280% 210 0.015 -0.01 167% 255 0.025 -0.012 148% 10 285 0.041 -0.01 124% 315 0.058 -0.011 1194 335 0.073 -0.009 1124 0.089 355 -0.007 108% 375 0.101 -0.006 106%

15

20

25

30

335

355

375

395

Escherichia coli 35218 Multiple Drug Resistance							
Time T=0	Control 0	105	* Inhib				
0	o	0					
60	0.001	-0.003	400%				
120	0.003	-0.003	200%				
180	0.008	-0.002	125%				
225	0.015	-0.003	1201				
260	0.026	0	100%				
285	0.033	0.002	941				
315	0.047	0.008	831				

0.062

0.075

0.085

0.101

Table 2N. Phospholipid Synthesis - Oligonucleotide #105

35

0.012

0.022

0.026

0.04

81%

71*

69%

Table 20. Periplasmic/Secretory Proteins - Oligonucleotide #46

	Escherichia coli 35218 Multiple Drug Resistance							
	Time T=0	Control 0.002	46	* Inhib				
5	0_	0	0					
	60	0.001	0.001	01				
	120	0.001	0.002	-1001				
l	180	0.001	0	100%				
	240	0.005	0.001	801				
	285	0.012	0.001	921				
.0	350	0.027	0.003	891				
	390	0.043	0.012	721				
	420	0.063	0.018	711				
	450	0.082	0.028	661				
	470	0.096	0.039	594				
5	500	0.106	0.046	571				

Table 2P. Transport Proteins - Oligonucleotide #114

	Salmonella typhimurium 23564							
20	Time T=0	Control 0.004	0.008	* Inhib				
,	0	0						
	60	-0.001	0					
	120	0	-0.002					
	165	0	-0.004					
25	230	0.003	-0.004	2331				
	260	0.005	-0.004	180%				
	305	0.014	-0.002	1141				
	335	0.021	0	100%				
İ	365	0.033	0.001	971				
İ	395	0.052	0.007	87%				
30	415	0.066	0.012	824				
	435	0.08	0.018	78%				
	455	0.093	0.026	72%				
L	476	0.108	0.035	68%				

	Table 2Q. Amino Acid Biosynthesis - Oligonucleotide #32							
	Escherichia coli 35218 Multiple Drug Resistance							
	Time T=0	Control 0	32 0.002	* Inhib				
5		0	0					
	60	0.002	0.001	50%				
	120	0.001	0	1001				
	180	0.003	0.001	671				
	240	0.008	0.001	884				
	285	0.015	0	100%				
10	320	0.026		100%				
Ì	350	0.04	0	100				
	380	0.058	0.002	971				
ı	410	0.076	0.002	97%				
	430	0.091	0.003	971				
15	450	0.105	0.003	971				
TO -				لاحصا				

	Table 2R. Lipo	polysaccharide	Synthesis -	Oligonuc	leotide #7		
	Escherichia coli 35218 Multiple Drug Resistance						
20	Time T=0	Control 0.006	73	0.005	* Inhib		
i				0			
	60	0		0_			
	120	0.001		0	100%		
	165	0.001		0	1001		
25	210	0.005		-0.001	120%		
•	240	0.008		0	100%		
	275	0.015		0	100%		
- 1	305	0.024		-0.001	104%		
1	335	0.034		0	100%		
30	365	0.048	-	0.001	981		
_	390	0.061		0.003	951		
į	410	0.07		0.003	961		
	430	0.086		0.005	944		
	455	0.1		0.01	901		

Table 2S. Purine/Pyrimidine Biosynthesis - Oligonucleotide #63 Escherichia coli 35218 Multiple Drug Resistance Control 0.002 Time T=0 t Inhib 0.004 0 0 0 5 60 0.001 0.001 01 120 0.001 0.002 -100% 180 0.001 0.001 01 240 0.005 0.002 60% 285 0.012 0.001 921 10 350 0.027 -0.001 104% 390 0.043 0.001 98% 420 0.063 0.002 97% 450 0.082 0.001 99 ¥ 470 0.096 0.004 961 500 0.106 0.008 15

Table 2T. Outer Membrane Proteins - Oligonucleotide #78

	Escherichia coli 15218 Multiple Drug Resistance						
20	Time T=0		Control 0.001	78 0.004	* Inhib		
		0	0	0			
		60	0.001	-0.002	300%		
	1	20	0.002	-0.002	200%		
ı	1	65	0.005	-0.003	160%		
25	2	10	0.015	-0.004	127%		
İ	2	\$5	0.025	-0.004	116%		
		85	0.041	-0.003	1074		
	3	15	0.058	-0.003	105%		
	3	35	0.073	-0.002	103%		
30	3	55	0.089	-0.002	102%		
, , , , , , , , , , , , , , , , , , ,	3	75	0.101	-0.002	102%		

	Table	2U. Nitrate Red	uctase - Oligonucleo	tide #71				
	Escherichia coli 35218 Multiple Drug Resistance							
	Time T=0	Control 0	71 0.002	* Inhib				
5	. 0		0					
	105	0.002	0	1001				
	175	0.003	-0.002	1678				
	220	0.004	-0.001	1251				
	270	0.007	-0.001	1148				
	300	0.012	-0.001	108%				
10	330	0.022	-0.001	105%				
	360	0.032	о	100%				
	395	0.052		100%				
	425	0.065	0.003	951				
	445	0.081	0.004	95%				
15	465	0.09	0.006	93 t				
	490	0.108	0.013	881				

	Table 2	V. Drug Resist	ance -	Oligonucleotide	#114		
20	Escherichia coli 15218 Multiple Drug Resistance						
	Time T=0	Control 0	114	0.006	† Inhib		
	0	0		0			
	105	0.002		-0.002	200%		
	175	0.003		-0.005	2671		
25	220	0.004		-0.003	1751		
	270	0.007		-0.003	1431		
	300	0.012		-0.004	1334		
ļ	330	0.022		-0.004	1181		
	360	0.032		-0.004	113%		
30	395	0.052		-0.004	108%		
	425	0.065		-0.003	105%		
	445	0.081		-0.001	101%		
	465	0.09		0	100%		
	490	0.108		0.004	961		

Table 2W. Vitamin Metabolism - Oligonucleoride #5

	Table 2W. Vitamin Metabolism - Oligonucleotide #5							
	Escherichia coli 35218 Multiple Drug Resistance							
	Time T=0	Control -0.001	5 0.002	* Inhib				
5	0	0	0					
٦	60	0.001	-0.001	2001				
	150	0.002	-0.003	250%				
٠	195	0.005	-0.002	140%				
	245	0.013	-0.001	108%				
	275	0.019	0	100%				
10	320	0.04	0	100%				
	350	0.054	-0.001	1024				
	365	0.066	0	100%				
	385	0.079	0	1001				
İ	415	0.095	-0.001	1011				
	430	0.105	0,001	991				
15 "		****						

Table 3A.

	Table JA.										
		Escherichia coli 15218 Multiple Drug Resistance									
	Time T=0	Control 0	A 11	nhib	В	0.006	Inhib		*Inhib		
20	0	0	0			0			0		
	60	0.001	-0.001	2001	-0.004		500%	-0.002	300%		
	105	0.002	-0.002	200%	-0.004		300%	-0.002	200%		
	145	0.002	-0.001	1501	-0.003		2501	-0.002	2001		
	190	0.002	-0.001	1501	-0.003		250*	-0.002	2001		
25	230	0.005	-0.001	1201	-0.003		160%	-0.002	140%		
	275	0.009	-0.001	1111	-0.003		1331	-0.003	1331		
	320	0.015	-0.002	1131	-0.002		1131	-0.003	120%		
	350	0.022	-0.001 1	1054	-0.001		1051	-0.003	1144		
	380	0.03	0	1001	-0.001		1034	-0.002	107%		
1	410	0.048	0.001	98%	-0.001		102%	-0.003	1064		
30	445	0.068	0.005	931	-0.003		1041	-0.003	104%		
	465	0.08	0.009	898	-0.002		1034	-0.003	1044		
į	485	0.097	0.015	854	0.002		984	-0.003	1031		

A=2'-0-Me version 18

B=12mer version 18

35 C=15mer version 18

Table 3B. Escherichia coli 35218 Multiple Drug Resistance 0.002 Control *Inhib Time T=0 5 0.001-0.001 2001-0.004 500¥ 0.002-0.001 105 1504 -0.004 3001 0.002-0.001 1501 -0.005 350% 190 0.002-0.001 150% -0.002 2004 230 0.005 -0.001 1201-0.004 1801 10 275 0.009-0.001 1111-0.004 1441 320 0.015-0.001 1078 -0.004 1271 350 0.022-0.001 1051-0.003 114 380 0.03-0.001 1031-0.003 110% 410 0.048-0.001 1021-0.003 106% 0.068-0.001 1011-0.001 1011 15 465 0.08-0.001 1011-0.001 101% 0.097-0.001 10110.002 485 981

D=5'amino group/15mer version 18

E=33mer version 18

20

						Tabl	e 3C.					
		Staphylococcus aureus 13301										
	Time T=0	Control 0	Α	0	*Inhib	В	*Inhib -0.001	c	*Inhil 0.001			
25	٥	0		0		0		0				
	90	0.003	0.002		331	0.003	01	0.003	01			
	150	0.003	0.001		671	0.004	-33%	0.003				
	210	0.005	0.002		601	0.004	20%	0.003	401			
	270	0.006	0.001		831	0.003	50%	0.003	50			
	325	0.014	0.001		934	0.002	86%	0.003	791			
30	380	0.032	0.002		941	0.003	911	0.002	941			
- 1	410	0.044	0.003		931	0.003	93%	0.003	931			
	440	0.057	0.004		931	0.003	95%	0.003	95%			
	470	0.075	0.005		931	0.021	72%	0.003	961			
	500	0.105	0.011		901	0.004	961	0.004	961			

35 A=2'-0-Me version 18

B=pEthoxy version 18

C=12mer version 18

				Tal	ole 3D	. `		
			Scaphyl	ococcu	s aure	us 1330	1	
	Time T=0	Control		0.003	*Inhib	E	0.003	*Inhib
	0			_ 0		<u> </u>	0	
5	65	0.001	0.001		01	0.003		-200%
	125	0.002	0.003		-501	0.003		-50%
	185	0.003	0.002		331	0.004		-331
	240	0.003	0.002		331	0.004		-334
	295	0.004	0.002		50%	0.003		25%
10	340	0.007	0.003		571	0.006		144
10	385	0.011	0.003		731	0.005		55%
i	415	0.016	0.002		88%	0.004	·	75%
	445	0.021	0.002		90%	0.004		811
	475	0.032	0.002		941	0.004	·····	884
	505	0.029	0.002		931	0.005		831
15	535	0.045	0.002		96%	0.006		871
	565	0.057	0.002		96%	0.005		911
	595	0.072	0.002		971	0.009		884
	625	0.09	0.002		984	0.006		931

D=15mer version 18

20 E=18mer version 18

25

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Table 4A. Salmonella typhimurium *Inhib39 0.004 %Inhib63 0.004 %Inhib 78 0.003 Control -0.001 18 *Inhib 0.002 5 90 0.001-0.001 200% -0.002 3001-0.002 3001-0.001 2001 150 0.002-0.004 300% -0.002 2008 -0.001 1501-0.001 1501 210 0.003-0.004 2331-0.002 1678 -0.001 1334 -0.001 1331 260 0.006-0.001 1174-0.001 1171-0.001 1174 -0.001 325 0.020 1004-0.001 1054-0.001 10510.001 95% 360 0.0330.002 9410.001 9710.002 9480.003 911 10 390 0.0490.007 8640.005 9010.004 9210.007 861 0.0670.012 420 8210.01 8510.007 90% 0.012 82% 0.0930.019 445 80% 0.016 8340.011 8810.019 801 460 0.1030.023 7810.02 8140.015 8510.024 771

15

		Table 4B.								
	Salmonella typhimurium									
20	Time T=0	Control 0.005		Inhib						
	0	0	0							
	60	-0.001	-0.001							
	120	0.001	-0.001	2001						
	165	0.003	-0.003	200%						
25	230	0.009	-0.004	1448						
	260	0.013	-0.004	1314						
	295	0.024	-0.003	1134						
	325	0.037	-0.002	105%						
]	350	0.051	-0.004	108%						
30	370	0.066	-0.003	105						
30	390	0.082	0	1001						
	410	0.098	-0.002	1024						
[430	0.112	0	1001						

Table 4C. Pseudomonas aeruginosa 0.005 Inhib 63 0.007 Control 39 0.002 0.005 0 0 5 0.001-0.001 20010 10010 100% 0.0020 1001-0.001 15010.002 0 % 250 0.003-0.001 1334-0.002 16710 100% 300 0.004-0.001 1251-0.002 15010 1001 345 0.0040 10010 10010 100% 375 0.0050.001 8010.001 8010.002 601 10 415 0.0080.003 6340.004 5010.004 501 0.0130.008 465 3810.007 4680.007 461 505 0.020.013 3510.013 3510.011 451 545 0.0360.022 3910.022 39%0.02 441 0.0510.038 2510.034 3310.034 334 0.0720.055 600 2410.052 2810.047

	(Table 4D.							
		Pseudomonas aeruginosa									
20	Time T≃0	Control C	_	*Inhi 0.004	114	0.002	Inhib				
				0		0					
	90	0.002	0.001	50	0.001		50%				
	120	0.003	0.002	331	0.003		01				
	180	0.006	0.003	501	0.004		331				
25	240	0.007	0.004	439	0.004		431				
	305	0.019	0.012	379	0.011		421				
	335	0.024	0.017	290	0.019		211				
	365	0.036	0.027	251	0.028		221				
	400	0.062	0.05	191	0.049		211				
30	420	0.074	0.061	189	0.06		194				
	440	0.086	0.074	148	0.071		171				
Į	460	0.103	0.091	12%	0.087		164				

Table 4E

					Table	72.					
	Klebsiella pneumoniae										
	Time T=0	Control 0.006		¥Inhib .008	78	0.006	*Inhib	-	*Inhib		
ļ	0	0	-0.001			0			0		
5	60	-0.002	-0.002		-0.001			-0.002			
	120	0	-0.003		-0.001			-0.0074			
i	165	0.004	-0.004	200%	-0.003		175%	-0.003	175%		
	230	0.011	-0.004	1364	-0.001		1091	-0.003	1274		
	260	0.019	-0.004	1214	0		100%	-0.003	116%		
10	295	0.036	-0.003	1081	0.003		921	-0.003	1084		
10	325	0.051	-0.001	102%	0.007		864	-0.003	1064		
	350	0.064	0	100%	0.012		81%	-0.003	105%		
İ	370	0.074	0.002	971	0.018		761	-0.003	1048		
·	390	0.088	0.006	934	0.025		721	-0.003	1031		
Į.	410	0.098	0.01	901	0.037		621	-0.003	1031		

15

20 Table 4F.

ļ	Klebsiella pneumoniae									
	Time T=0	Control 0.006		0.009	Inhib	111	0.008	*Inhib		
	0	0		0			0			
	60	-0.001	-0.00	3		-0.002				
25	135	0.005	0		100%	-0.002		2001		
	180	0.012	0		1001	0		100%		
	210	0.019	0.004		791	0.002		834		
- 1	240	0.03	0.006		801	0.006		671		
	270	0.05	0.014		72 %	0.012		631		
30	315	0.072	0.03		58%	0.024		541		
30	335	0.083	0.039		534	0.032		541		
	355	0.107	0.051		521	0.041		511		

Table 4G. Yersinia mollaretti *Inhib4 \$Inhib127 Control 2 0.003 *Inhib 5 010.001 0.0010.001 010.001 0 % 0.0020.002 010.002 010.002 01 2510.003 0.0040.003 2510.003 25% 0.0080.003 6310.003 6310.004 501 0.010.004 6010.004 6010.006 401 10 0.0140.008 4310.008 4310.012 14% 350 0.0230.012 4810.013 4310.018 221 380 0.0290.018 3810.018 3840.025 14% 0.0390.026 3310.027 3110.035 101 0.0540.035 3510.036 3310.048 11% 0.0750.05 3310.056 2510.071 51 15 2610.087 500 0.0960.07 2710.071 9 %

2910.075

2610.092

20		<u> </u>		Table 4H.	·							
	<u></u>	Yersinia mollaretti										
	Time T=0	Control 0.002		*Inhib 0.004	73	0.004	lnhib					
		0		0		0						
	90	0.001	0.002	-1001	o		100%					
25	190	0.002	0.003	-501	0.001		50%					
	250	0.003	0.003	01	0.001		671					
	300	0.003	D	100	0.001		671					
	345	0.006	0.003	501	0.003		50%					
	375	0.008	0.005	381	0.005		38%					
	415	0.013	0.008	381	0.009		314					
30	465	0.023	0.018	221	0.019		178					
	505	0.031	0.027	131	0.027		18%					
	545	0.055	0.043	221	0.043		221					
	575	0.074	0.065	121	0.064		148					
ļ	605	0.093	0.083	114	0.08		141					
35	615	0.103	0.089	144	0.088		15%					

0.1010.072

				Tab.	le 41.					
				Neiss	ria si	cca				
	Time T=0	Pos. Control OD	16	*Inhib	12		*Inhib	20		*Inhib
		0.029	0.06	4		0.035			0.084	
5	°	•				0			0	
•	30	0.002	-0.003		-0.002			-0.002		
ļ	65	0.002	-0.003		-0.003			-0.004		
	125	0.006	-0.001	1174	-0.002		1334	-0.002		133%
	150	0.01	0.001	901	a		100%	-0.002		120%
10	180	0.014	0.001	934	0.002		864	-0.001		107%
10	240	0.023	0	1001	0.002		914	-0.003		1134
	300	0.029	0.01	661	0.009		691	0.006		791
	330	0.029	0.014	521	0.013		551	0.012		59%
	390	0.033	0.014	581	0.009		73%	0.012		641
	450	0.031	0.004	874	0.009		711	0.003		90%
15	490	0.036	0.014	61%	3.008		781	0.008		78%
1	520	0.038	0.015	619	0.014		631	0.011	******	718
	560	0.049	0.013	73%	0.002			0.007		86%
	590	0.052	0.017	678	0.014		731	0.012		778
1	620	0.057	0.018	681	0.014		751	0.014		75%
	650	0.059	0.016	7310	.018			0.014		76%
20	680	0.063	0.018	7110	.016			0.016		75%
	710	0.068	.019		.017			0.016		76%

25

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Table 4J. Neisseria sicca Pos. Control OD 0.029 Time T=0 *Inhib 15 *Inhib 5 0.018 0.056 ٥ 30 0.002-0.001 0.001 0.002-0.004 65 125 0.006-0.001 11780.002 67% 150 0.010.004 601-0.003 1301 10 180 0.0140.005 641-0.002 1141 240 0.0250.004 844-0.003 112* 7010.01 300 0.0270.008 634 330 0.0290.015 4810.018 381 390 0.0330.012 644 -0.003 1091 15 450 0.0310.005 8410.01 681 490 0.0360.012 6710.016 561 0.0380.007 520 8210.018 531 560 0.0490.011 78 0 . 021 571 590 0.0520.011 7910.02 621 620 0.0570.011 8110.018 68% 20 0.0590.012 650 8010.018 691 680 0.0630.011 8310.02 681 0.0680.012 8210.17 710 75%

				Tal	ble 4K	·				
25				Serrat	ia liq	uefacien	s			
	Time T=0	Control -0.001	2 -0	%Inhib .001		¥Inhib 001		*Inhib 002	114	*Inhib
	٥	0		0		0		0		0
	110	0.002	0.002	0%	0.002	04	0	100%	0.002	0 %
	180	0.003	0.003	0%	0.001	671	0.001	671	0.002	331
30	240	0.003	0.002	33%	0.001	671	0.001	671	0.002	331
	300	0.002	0.002	0.8	0.001	50%	0	100%	0.001	50%
	360	0.005	0.002	60%	0.001	80%	0	. 100%	0.001	80%
	420	0.011	0.003	73%	0.001	914	0.001	911	0.002	821
į	475	0.022	0.003	861	0.002	918	0.001	951	0.003	86%
35	520	0.041	0.003	93%	0.001	98%	0.001	981	0.002	95%
35	610	0.082	0.003	964	0.001	991	0.001	991	0.002	984
	655	0.1	0.003	971	0.001	994	0.001	991	0.002	981

	Table 4L.									
			s	reptococ	cus mu	cans				
	Time T=0	Control 0.184	1 0.	*Inhib 187		*Inhil		*Inhib		
		0		0		0		0		
5	60	0.001	-0.003	400%	-0.001	2001	-0.002	300%		
:	115	0.006	-0.001	1178	0.003	501	0.001	81%		
	145	0.011	-0.001	1094	0.003	731	0.003	73%		
	180	0.016	0.002	881	0.008	50%	0.006	63%		
	210	0.022	0.004	824	0.01	551	0.008	64%		
10	245	0.031	0.009	718	0.015	521	0.014	554		
10	290	0.047	0.015	68%	0.021	551	0.021	55%		
- 1	320	0.059	0.022	63%	0.026	561	0.03	49%		
į	350	0.071	0.03	581	0.032	551	0.04	445		
l	385	0.082	0.036	56%	0.032	611	0.047	431		
l	415	0.097	0.042	578	0.036	631	0.05	481		
15	445	0.109	0.045	594	0.039	641	0.063	42%		

				Table 4M.	7	
		S	treptoc	occus muta	ns	
20	Time T=0	Control 0.184	132	*Inhib .187		ŧInhib .183
	0			0		0
	60	0.001	-0.002	300%	-0.003	400%
	115	0.006	0.001	834	-0.001	1178
	145	0.011	0.001	911	0.002	821
25	180	0.016	0.006	634	0.004	75%
	210	0.022	0.008	641	0.008	648
	245	0.031	0.01	681	0.013	581
	290	0.047	0.017	644	0.025	478
	320	0.059	0.022	638	0.034	421
30	350	0.071	0.027	621	0.045	371
	385	0.082	0.028	664	0.054	341
	415	0.097	0.033	661	0.062	361
į	445	0.109	0.034	691	0.069	371

Table 4N. Streptococcus pyogenes †Inhib 0.179 *Inhib 89 0.179 Control 0.177 Time T=0 5 0.0010 1001-0.001 2001-0.004 500% 170 0.003 -0.002 1678 -0.002 1674 -0.005 2671 210 0.005-0.001 12010 1001-0.003 1604 240 0.008-0.001 1134 -0.001 1131-0.002 125% 300 0.010 10010.001 9010 100% 345 0.0140.003 7910.002 8610 100% 10 390 0.0210.006 7110.003 8610 100% 0.0360.01 450 7210.008 7810.007 814 0.0670.017 510 7510.015 7810.015 781 0.0930.025

73 \$ 0.026

7410.029

7210.025

7310.025

73%

773

15

540

555

0.1070.028

			Tab	le 40.				
20		s	trepto	occus	pyoger	ies		
	Time T=0	Control 0.177	132	0.177	*Inhib	114	0.181	*Inhib
	0	0		0			a	
	110	0.001	-0.001		200%	-0.001		1334
25	170	0.003	-0.003		200%	-0.003		1751
	210	0.005	0		100%	-0.004		2001
	240	0.008	-0.001		1134	-0.001		1174
	300	0.01	0.001		901	0		100%
	345	0.014	0.002		86%	0.001		911
	390	0.021	0.004		814	0.005		691
ļ	450	0.036	0.009		75%	0.015		55%
30	510	0.067	0.015		78%	0.031		478
	540	0.093	0.021		77%	0.047		451
	555	0.107	0.021		80%	0.053		481

Table 4P. Shigella 0.003 tInh Control 0.001 Time T=0 **†Inh** *Inh 0.004 127 0.003 0 5 95 0.001 -0.001 2001 -0.001 200% -0.001 2001 155 0.005 -0.001 120% -0.003 1604 -0.002 140% 215 0.009 -0.001 1111 -0.002 1224 -0.002 1221 275 0.027 100% -0.002 107% -0.001 104% 305 0.038 100% -0.003 108% -0.002 105% 335 0.044 0.001 981 -0.001 102% -0.003 107% 10 365 0.047 0.004 914 -0.002 104% -0.001 102% 395 0.051 889 -0.002 1041 -0.001 102% 425 0.051 0.008 84% -0.003 106% -0.001 102%

15

20			Table 40.			
20			Shigella			
	Time T=0	Control 0.001	132	*Inh	114	tInh 03
	0	0	0	o		
25	95	0.001	-0.001	2001	-0.001	200%
	155	0.005	-0.001	120%	-0.002	140%
	215	0.009	-0.001	1111	-0.003	1331
- 1	275	0.027	-0.001	104%	-0.003	111*
	305	0.038	-0.002	105%	-0.003	108%
	335	0.044	-0.003	107%	-0.003	107%
	365	0.047	-0.001	102*	-0.003_	1064
30	395	0.051	0	100%	-0.002	104%
	425	0.051	0	100%	-0.002	104%

	<u> </u>	Table 4R.										
		Наеторһ	ilus									
	Time T=0	Control 0.161	78	*Inh 0.017								
		0		0								
5	70	0.007	0	100%								
	140	0.012	0.008	334								
	190	0.013	0.01	234								
ı	235	0.013	0.013	0%								
- 1	275	0.013	0.013	01								
10	305	0.015	0.012	20%								
10	365	0.016	0.013	19%								
	24'	0.026	0.011	581								
	29. 20.	0.051	0.014	73%								
	46'	0.241	0.021	91%								

			Table 45	S			
		м	ycobacte	rium			
Time T=0	Control 0.167	114	*Inh	10	¥Inh 68	21 0.1	*Inh
	0					0	
90	0.006	0.001	834	0.001	831	0.002	67%
120	0.009	0.003	671	0.002	78%	0.006	331
165	0.014	0.005	648	0.005	641	0.01	29%
195	0.021	0.006	71%	0.005	761	0.008	624
240	0.021	0.007	671	0.007	671	0.009	571
270	0.018	0.013	28%	0.01	441	0.013	281
305	0.028	0.016	431	0.012	571	0.014	501
405	0.04	0.026	35%	0.032	201	0.025	381
465	0.051	0.032	37%	0.041	201	0.032	371
525	0.063	0.04	371	0.051	191	0.043	324
555	0.073	0.046	371	0.06	184	0.052	291
585	0.08	0.051	361	0.065	194	0.055	314
615	0.085	0.062	271	0.073	141	0.062	27%
645	0.097	0.065	331	0.079	191	0.068	304
	7=0 0 90 120 165 195 240 270 305 405 465 525 555 585 615	T=0 0.167 0 0 90 0.006 120 0.009 165 0.014 195 0.021 240 0.021 270 0.018 305 0.028 405 0.04 465 0.051 525 0.063 555 0.073 585 0.08 615 0.085	Time Control 0.167 0. 0 0 0 0 0 90 0.006 0.001 120 0.009 0.003 165 0.014 0.005 195 0.021 0.006 240 0.021 0.007 270 0.018 0.013 305 0.028 0.016 405 0.04 0.026 465 0.051 0.032 525 0.063 0.04 555 0.073 0.046 585 0.08 0.051	Time Control 114 0.163 0 0 0 90 0.006 0.001 83% 120 0.009 0.003 67% 165 0.014 0.005 64% 195 0.021 0.006 71% 240 0.021 0.007 67% 270 0.018 0.013 28% 305 0.028 0.016 43% 405 0.04 0.026 35% 465 0.051 0.032 37% 525 0.063 0.04 37% 555 0.073 0.046 37% 585 0.08 0.051 36% 615 0.085 0.062 27%	T=0 0.167 0.163 0.1 0 0 0 0 90 0.006 0.001 83% 0.001 120 0.009 0.003 67% 0.002 165 0.014 0.005 64% 0.005 195 0.021 0.006 71% 0.005 240 0.021 0.007 67% 0.007 270 0.018 0.013 28% 0.01 305 0.028 0.016 43% 0.012 405 0.04 0.026 35% 0.032 465 0.051 0.032 37% 0.041 525 0.063 0.04 37% 0.06 585 0.073 0.046 37% 0.06 585 0.08 0.051 36% 0.065 615 0.085 0.062 27% 0.073	Mycobacterium Time T=0 Control 0.167 114 0.163 \$Inh 0.168 10 0.168 0 0 0 0 0 0 90 0.006 0.001 83\$ 0.001 83\$ 120 0.009 0.003 67\$ 0.002 78\$ 165 0.014 0.005 64\$ 0.005 64\$ 195 0.021 0.006 71\$ 0.005 76\$ 240 0.021 0.007 67\$ 0.007 67\$ 270 0.018 0.013 28\$ 0.01 44\$ 305 0.028 0.016 43\$ 0.012 57\$ 405 0.04 0.026 35\$ 0.032 20\$ 465 0.051 0.032 37\$ 0.041 20\$ 525 0.063 0.04 37\$ 0.061 18\$ 585 0.08 0.051 36\$ 0.065 19\$ 615	Mycobacterium Time T=0 Control O.167 114 O.163 VInh O.168 21 O.168 0.002 0.002 0.002 0.006 0.006 0.003 674 0.002 783 0.006 0.006 0.006 0.005 644 0.001 0.006 0.006 0.005 644 0.001 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.009

			Table 4T			
		M)	cobacter:	i um		
	Time T=0	Control 0.167	18 0.1	¥Inh 63	78 *In:	
		0	0			,
5	90	0.006	-0.001	_1170	0	100%
	120	0.009	0.002	781	0.003	671
	165	0.014	0.007	501	0.003	791
	195	0.021	0.006	714	0.004	814
	240	0.021	0.008	621	0.006	714
10	270	0.018	0.008	56%	0.003	831
10	305	0.028	0.01	644	0.009	684
	405	0.04	0.022	451	0.018	55%
	465	0.051	0.03	42%	0.024	531
	525	0.063	0.037	418	0.029	541
	555	0.073	0.044	40%	0.037	498
15	585	0.08	0.047	411	0.04	50%
	615	0.085	0.052	394	0.042	511
<u>[</u>	645	0.097	0.059	39%	0.056	424

		Table 4U.										
		Helicoba	ccer									
	Time T=0	Control 0.08	78	*Inh								
	0	0		0								
25	70	-0.004	-0.009									
	140	0	-0.006									
	190	0.001	-0.005	6001								
ſ	235	0.003	-0.001	1331								
l	275	0.004	0	1001								
	305	0.009	0.004	56%								
30	365	0.01	0.003	70%								
	24'	0.057	0.01	821								
	29' 50'	0.065	0.012	82%								
	46'	0.065	0.005	921								

Table 4V. Enterococcus Time T=0 Control 0.09 127 %Inh 0.087 *Inh 0.088 *Inh 0.088 7 %Inh 0.086 p127 0 0 5 60 0 -0.004 -0.006 -0.006 -0.007 105 0.005 -0.004 1804 -0.002 1401 -0.003 160% -0.005 2001 150 0.026 0.008 0.009 0.008 691 0.01 624 170 0.066 0.029 56% 0.029 561 0.025 62% 0.032 521 195 0.076 0.04 478 0.04 47% 0.036 53**%** 0.043 431 210 0.091 0.051 441 0.052 431 0.047 10 48% 0.054 411 215 0.11 0.062 0.064 421 0.055 · 50% 0.066 40%

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	<u></u>		Tab	le 4W.		
			Enteroc	occus		
	Time T=0	Control 0.042	0.0	linh	76 %Ir	
	0	, o	0		0	
20	60	0.002	-0.002	200%	-0.001	150%
	120	0.006	-0.001	1171	0	1001
	160	0.023	0.002	911	0.003	871
	190	0.036	0.01	721	0.013	64%
	210	0.051	0.015	718	0.02	618
	230	0.074	0.031	581	0.04	461
25	245	0.083	0.037	551	0.046	45%
	255	0.094	0.047	50%	0.057	391
Į	265	0.109	0.054	501	0.065	401

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	<u></u>	·	Table 4X.					
		,	Streptoc	occus p	neumonia			
	Time T=0	Control 0.17	0.1	%Inh 72	78 0.1	*Inh .74	114	*Inh 7
_							0	
5	60	0.004	0	100%	0	100%	-0.001	125%
	110	0.003	-0.005	2671	-0.001	1334	-0.001	1331
	170	0.003	-0.003	2001	-0.001	1334	-0.001	1334
	220	0.004	-0.002	1501	0	100%	-0.001	125%
	260	0.004	-0.001	1251	-0.001	1251	-0.001	1251
10	310	0.007	-0.002	1291	0	1001	-0.001	1148
10	370	0.008	-0.003	1384	0	100%	0	100%
	445	0.009	-0.002	1221	0	100%	0	100%
	485	0.009	-0.003	1331	0.001	891	0.001	891
	19,32,	0.014	0.001	931	0.011	211	0.008	43%
	21.12.	0.014	0.001	931	0.01	291	0.006	571
15	23,32,	0.015	0.002	871	0.012	20%	0.008	471
	27'	0.016	0.001	941	0.013	19%	0.009	441
	28'30'	0.016	0.002	881	0.014	121	0.01	381
	45'20'	0.023	0.018	221	0.024	-41	0.018	22%
ļ	48'20'	0.024	0.008	671	0.025	-48	0.014	42%
20	51.50.	0.024	0.01	584	0.035	-461	0.022	8 %
20	54'20'	0.026	0.011	58%	0.028	-8%	0.021	19%
	70'35'	0.035	0.014	60%	0.033	61	0.027	23%
	95'35'	0.05	0.025	501	0.059	-181	0.04	201
į	101.	0.068	0.025	631	0.046	321	0.043	37%

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	(7	able 4Y.			
		Strepto	ococcus pr	neumoni	a	
	Time T=0	Control 0.17	127	127 *Inh 0.172		% Inh
)
5	60	0.004	-0.001	1251	-0.001	125%
	110	0.003	-0.001	1331	-0.003	2001
	170	0.003	-0.002	1671	-0.003	200%
	220	0.004	-0.002	150%	-0.002	150%
	260	0.004	-0.001	125%	-0.002	150%
10	310	0.007	-0.002	1291	-0.001	114*
10	370	0.008	o	100%	0	100%
10	445	0.009	0	100%	0	100%
	485	0.009	0	1001	0	100%
	19'35'	0.014	0.008	431	0.009	36%
	21'35'	0.014	0.007	501	0.009	36%
15	23'35'	0.015	0.008	471	0.009	401
	27'	0.016	0.01	378	0.013	19%
	28,30,	0.016	0.012	254	0.012	25
	45'20'	0.023	0.019	178	0.022	41
	48'20'	0.024	0.2	17%	0.021	134
20	51.50.	0.024	0.021	124	0.022	81
20	54'20'	0.026	0.022	151	0.024	81
j	70'35'	0.035	0.027	234	0.033	61
į	95'35'	0.05	0.048	41	0.05	01
	101.	0.068	0.048	291	0.052	241

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Table 4Z

		Iddie 42.						
	<u></u>		Vi	brio				
	Time . T=0	Control 0.138	78 0.1	*Inh	127	*Inh		
	0	0	0		0			
5	. 70	0.002	-0.001	150%	-0.003	2501		
	140	0.002	0	100%	-0.002	200%		
	190	0.005	0	100%	0	100%		
	235	0.005	0.001	80%	-0.002	140%		
	275	0.005	0.001	80%	-0.003	160		
7.0	305	0.005	0	100%	0	100%		
10	365	0.004	-0.001	125%	-0.002	150%		
	24'	0.006	0.003	50%	0	100%		
	46'	0.177	0.006	97%	0.129	27%		

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Tạble SA. Staphylococcus aureus 13301 0.002 Inhib85 Control 0.001 *Inhib *Inhib|68 0 5 65 0.0010.001 010.002 -10010.001 01 125 0.0020.002 010.003 -5010.002 Oŧ 185 0.0030.002 3340.003 010.003 01 240 0.0030.002 33 0 . 003 010.002 334 295 0.0040.001 7510.003 2510.002 501 340 0.0070.002 7110.003 5710.003 571 10 385 0.0110.004 6410.003 7310.002 821 415 0.0160.002 8810.003 8110.001 941 0.0210.002 445 9010.003 8610.002 90 **t** 475 0.0320.002 9410.003 9110.002 941 505 0.0290.001 9710.003 9010.002 93% 15 535 0.045 0.001 9810.003 9310.002 961 565 0.0570 10010.001 9810.003 95% 0.0720.002 595 9710.003 9610.003 961 625 0.090.002 9810.002 9810.002 981 0.456-0.002 10010 10010.026 941

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Table 58

	Table 58.							
		Stapi	yloco	cus aureus	13301			
	Time T=0	Control 0.00	112	*Inhib 0.005	18	*Inhib 0.003		
	0			0	<u> </u>	0		
5	55	0.001	0	1001	0.001	01		
	125	0.002	0.002	01	0.003	-50%		
	185	0.003	0.001	671	0.002	331		
	240	0.003	0.001	67%	0.002	331		
	295	0.004	0.002	50%	0.002	50%		
10	340	0.007	0.001	861	0.003	571		
10	385	0.011	0.001	914	0.003	731		
	415	0.016	0	100%	0.002	881		
	445	0.021	0	100%	0.002	901		
	475	0.032	0.001	97%	0.002	941		
	505	0.029	0.001	971	0.002	934		
15	535	0.045	0.002	96%	0.002	961		
	563	0.057	0.002	96%	0.002	96%		
	5 9 5	0.072	0.001	991	0.002	97%		
	625	0.09	0	1001	0.002	984		
į	25.	0.456	-0.003	1011	0	100%		

					Table	5C.				
			ŀ			coli i g Resis				
25	Time T≃0	Control 0.001	21	0.004	lnhib	68	0.005	*Inhib		*Inhib
		0		0		<u> </u>	0.001			0.001
	70	0.002	0.001		50%	-0.002		200%	-0.001	150%
	130	0.002	0.001		50%	-0.001		150%	-0.001	1501
	190	0.002	-0.001		150%	-0.003		250%	-0.002	2001
	250	0.009	-0.002		1221	-0.003		133%	-0.003	1331
	295	0.015	-0.002		1134	-0.002		1134	-0.002	113%
30	325	0.024	-0.001		104%	-0.002		1084	-0.002	1081
	355	0.032	-0.002		106%	-0.002		1061	-0.002	1061
	385	0.046	-0.002		1041	-0.003		107%	-0.002	104%
	415	0.068	-0.001		1011	-0.002		1034	-0.002	1031
	445	0.087	-0.001		101%	-0.001		101%	-0.001	1014
35	465	0.1	-0.001		1011	-0.001		1014	-0.002	1021
	\$55	0.138	0.009		931	0.01		931	0.005	961
	27'	0.191	0.196		-38	0.192		-11	0.192	-11

Escherichia coli 35218 Multiple Drug Resistance 0.004 Inhib18 Control 112 0.001 Time T=0 *Inhib -0.002 5 70 0.002-0.004 3001-0.001 150% 0.002-0.005 130 350%-0.001 150% 190 0.002-0.005 3501-0.001 150¥ 250 0.009-0.005 1561-0.001 1111 295 0.015-0.004 12710 1001 10 325 0.024-0.004 1178-0.001 104 355 0.032-0.005 11640 100% 385 0.046-0.004 1091-0.001 1024 415 0.068-0.004 10610 1001 445 0.087-0.003 10310.003 971 465 0.1-0.004 10410.004 961 15 555 0.1380.008 9410.026 811 0.1910.178 710.174

20	Table 5E.							
		NBT89			oli 25922 ent concer		ions	
	Time 7.0	Control	3.1mg 0.004) Int	1.05mg 0.003) Ir.	0.525mg 0.00	, \ inhib
		с		_			3	
25	50	0.001	-0.001	2001	-0.001	200	-0.001	3003
	:2:	0.001	-0.002	1001	-0.001	2001	-0.001	2001
	225	0.005	-0.002	1404	-0.001	1201	-0.001	1201
	275	0.012	-0.001	1081	-0.001	1081	-0.001	1064
	315	0.027	-0.001	1041	-0.001	1041	0	1001
	375	0.035	-0.001	1031	-0.001	1034	0.001	978
	355	0.044	-0.002	1051	-0.001	1021	0.002	951
	375	0.052	0.002	1049	-0.001	1071	0.002	963
30	395	9.06	0.003	1051	-0.001	1021	0.002	979
	415	0.081	0.002	1021	-0.001	1011	0.00)	961
	410	0.092	0.002	1021	0.001	1011	0.005	951
	445	0.101	0.002	1021	.000	1001	0.009	911
1	24 NE			271		181		161

	-				Table 5	<u>r.</u>				
	<u></u>	Escheric	hia coli 2592	2	WBT49	At dif	ferent con	Centrat	ion s	
	Time T-0	Control	0.265mg 11n 0.003	hib	0.133mg 0	tinhi	3.07mg 0.03) lahib	. 0 3 5mg	*Inhib
				_						3
_	60	0.001	-0.001 2	001	0	2031	9	::24	0	1001
5	120	0.001	-0.001 2	001	0	:001	-0.001	::01		1001
	225	0.005	-0.001 1	201	0	1001	9.001		0.002	601
	270	0.012	-0.001 10	18 t	0.603	751	0.004		0.006	501
	315	0.027	0.003	91	0.01	931	0.012	531	0.015	441
	335	0.035	0.004	,,,	0.015	571	0.018	431	0.022	373
	155	0.044	0.006	64	0.021	521	0.024	450	7.029	349
10	375	0.052	.008 6	51	0.025	521	0.029	449	0.035	333
	795	0.06	0.012	01	0.032	474	0.037	3310	.044	279
	415	0.001	.016 7	83	0.044	461	0.052	3630	.061	253
	430	0.092	.021 7	71	3.054	4:1	0.063	3290	.072	221
	445	0.101	.028 7	210	0.064	371	7.073	211 2	.092	193
	24 hr		. 1	••		::)		:51		111

Table 6A

The Effects of Oligonucleotide Purification Method on the Percent Inhibition of Escherichia coli 35218 (See Section 5.5.)

			1366.3	ection 5.	5.}		
	Time	Control	À	В	С	D	ε
	0	0					
5	90	.003	100%	100%	100%	1001	1001
	150	.004	100%	100%	100%	100%	100%
	220	.008	75%	100%	1001	63 t	100%
	270	.014	364	1001	1001	141	100%
	315	.029	381	1001	100%	104	1001
10	345	.038	218	1001	100%	81	100%
10	375	.059	251	931	971	31	100%
	400	.079	271	901	901	64	994
	420	.089	25%	841	841	5%	981
	435	.099	241	831	834	61	96%

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Table 6B
The Effects of Oligonucleotide Purification Method on the Percent Inhibition of Escherichia coli 15218 (See Section 5.5.)

	Time	Control	F	G	н	1
20	0	0				
	90	.003	1001	1001	100%	100%
	150	.004	100%	100%	1001	1001
	220	.008	100%	100%	100%	100%
	270	.014	1001	100%	1001	100%
	315	.029	631	100%	100%	1001
25	345	.038	478	100%	1001	100%
	375	.059	504	100%	981	1001
	400	. 079	341	96%	911	100∜
ł	420	.089	431	961	88%	1001
[435	.099	418	931	861	1001

Table 7
Antigene Oligonucleotides Targeted to DNA sense strand for Triplex Formation

				richia le Drug				
_	Time T=0	Control 0.002	96.SS	0.008	*Inhib	73.SS	0.004	*Inhib
5	۰	0		,0.004			0	
	60	0.001	0		1001	0.001		01
	120	0.001	0		100%	0		100%
	180	0.001	0	 -	100%	-0.001		2001
	240	0.005	-0.001		1204	0		1001
10	285	0.012	-0.001		1081	-0.002		1171
İ	350	0.027	-0.001		104%	0		1001
- }	390	0.043	0.002		95%	0.001		981
	420	0.063	0.006		90%	0.004		941
i	450	0.082			881	0.008		90%
15	470	0.096			824	0.01		90%
[500	0.106	0.023		78%	0.012		891

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Table 8A. Escherichia coli 11370 Streptomycin Resistant Control 73 *Inhib 0.004 0 0 0 5 60 0.005 0 100% 140 0.011 -0.002 118% 170 0.013 100% 0 215 0.021 0.003 86% 245 0.032 0.005 841 10 275 0.045 0.007 841 305 0.062 0.009 85% 325 0.076 0.009 88% 340 0.09 0.01 891 350 0.1 0.012 88%

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Table 8B Escherichia coli 29214 Sulfonamide Resistant 20 Time T=0 Control 0.001 73 *Inhib 0.003 0 0 50 0.001 -0.002 300% 130 0.005 -0.001 120% 25 175 0.015 -0.001 1071 205 0.022 -0.001 105% 235 0.031 -0.001 1034 270 0.05 0 100% 295 0.065 0 100% 315 0.081 0.003 96% 30 335 0.092 0.006 931

Table 8C

	Table 8C.								
	Int	Escherichia coli 25922 Intermediate Penicillin Resistant							
	Time T=0	Control 0.004	73 0.006	*Inhib					
5	0	0	0						
	60	-0.00 1	-0.001						
	120	0	-0.001						
	165	0	-0.001						
	230	0.003	-0.001	1331					
	260	0.005	-0.002	140%					
10	305	0.014	-0.002	114%					
	335	0.021	-0.002	110%					
	365	0.033	-0.002	106%					
	395	0.052	-0.001	1021					
	415	0.066	-0.002	103%					
15	435	0.08	-0.002	103%					
	455	0.093	-0.002	1024					
Į	475	0.108	-0.001	1014					

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able SD

		Salmone.	lla typhimurium 23564	
	Time T=0	Control 0.005	73 0.00s	tInhib
	0	0	0	
25	60	0.001	-0.001	
	120	0.001	-0.001	2001
	165	0.003	-0.003	200%
	230	0.009	-0.004	1448
	260	0.013	-0.004	1314
	295	0.024	-0.003	1134
30	325	0.037	-0.002	105%
	350	0.051	-0.004	108%
	370	0.066	-0.003	105
	390	0.082	0	100%
	410	0.098	-0.002	1023

Table 8E

		Klebsiel.	la pneumoniae 4352				
	Time T=0	Control 0.006	73	*Inhib			
	o	0	0				
5	60	-0.00 2	-0.002				
•	120	0	-0.0074				
	165	0.004	-0.003	1751			
	230	0.011	-0.003	127%			
	260	0.019	-0.003	116%			
10	295	0.036	-0.003	108%			
j	325	0.051	-0.003	106%			
	350	0.064	-0.003	1051			
	370	0.074	-0.003	104%			
į,	390	0.088	-0.003	103%			
, [410	0.098	-0.003	103%			

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20	F	Table 8F.								
20		Escherichia coli 15218 Multiple Drug Resistance								
	Time T=0		Control 0.001	73 -0.004	*Inhib					
		o	0	0						
	<u></u>	60	0.001	-0.003	400%					
25		120	0.003	-0.002	1671					
		180	0.013	-0.001	1081					
		210	0.019	-0.002	1114					
		240	0.027	-0.001	1041					
		270	0.04	0	100%					
1		300	0.058	0.003	951					
30		320	0.075	0.006	921					
		340	0.089	0.008	91%					
ł		355	0.103	0.013	87%					

Table 8	IG
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	Table 30.						
	<u></u>						
	Time T=0	Control 0	73 -0.007	*Inhib			
		0	0				
5	60	0	-0.003				
	120	0.003	-0.004	2334			
	165	0.006	-0.003	1504			
	210	0.01	0.001	90%			
:	240	0.014	0.004	71%			
10	270	0.024	0.011	541			
10	300	0.034	0.021	38%			
	340	0.48	0.033	314			
	360	0.06	0.041	32%			
	380	0.072	0.05	31%			
	400	0.09	0.062	31%			
15	420	0.102	0.07	314			

25

30

Tables 9(A-G)
Oligonucleotide NBT 114 vs. Different Strains of Bacteria

5 Table 9A Escherichia coli 11370 Streptomycin Resistant Time T=0 Control 114 * Inhib 0.004 0 0 o 60 0.005 -0.003 160% 10 140 0.011 0 100% 170 0.013 0.003 778 215 0.021 0.009 571 245 0.032 0.014 561 275 0.045 0.018 60% 15 305 0.062 0.024 61% 325 0.076 0.03 61% 340 0.09 0.034 621 350 0.1 0.036

20

	Table 9B.								
	Escherichia coli 29214 Sulfonamide Resistant								
	Time T=0		Control 0.001	114 t	Inhib				
٦-		0	0	0					
25		60	0.001	-0.002	3001				
		130	0.005	-0.001	1201				
		175	0.015	-0.001	1071				
		205	0.022	-0.001	1051				
1		235	0.031	0	1001				
30		270	0.05	0.005	901				
		295	0.065	0.007	891				
ı		315	0.081	0.012	85%				
		335	0.092	0.017	821				

Table 9C.

	Table 90.						
	Escherichia coli 25922 Intermediate Penicillin Resistant						
	Time T=0		Control 0.004	0.008	Inhib		
_		0	0	o			
5		60	-0.001	0			
		120	0	-0.002	•		
		165	0	-0.004			
		230	0.003	-0.004	2331		
		260	0.005	-0.004	1804		
10		305	0.014	-0.002	1148		
		335	0.021	00	100%		
		365	0.033	0.001	978		
		395	0.052	0.007	87%		
		415	0.066	0.012	821		
		435	0.08	0.018	781		
15	ļ	455	0.093	0.026	721		
	<u></u>	475	0.108	0.035	88%		

Table 9D.

20	Salmonella typhimurium 23564						
	Time T=0	Control 0.005	0.007	Inhib			
		0	0				
25	60	-0.001	0				
	120	0.001	-0.001	2001			
	165	0.003	-0.003	200%			
	230	0.009	-0.003	1331			
l	260	0.013	-0.002	115%			
	295	0.024	0	1004			
	325	0.037	0.003	921			
30	350	0.051	0.009	824			
	370	0.066	0.012	821			
İ	390	0.082	0.017	791			
Į	410	0.098	0.024	764			

Table Of

	Table 9E.						
	Klebsiella pneumoniae 4352						
	Time T=0		Control 0.006	114	0.008	Inhib	
		٥			-0.001		
5		60	-0.002		-0.002		
		120	0		-0.003		
		165	0.004		-0.004	2001	
- 1		230	0.011		-0.004	136%	
		260	0.019		-0.004	1214	
,,		295	0.036		-0.003	108%	
10		325	0.051		-0.001	1021	
		150	0.064		0	100%	
		170	0.074		0.002	978	
1	3	90	0.088		0.006	931	
L	4	10	0.098		0.01	901	

Table 9

	Escherichia coli 15218 Multiple Drug Resistance							
20	Time T=0	Control 0.001	0.003	Inhib				
		0						
	60	0.001	-0.002	3001				
	120.	0.003	-0.001	1331				
	180	0.013	· . 0	100%				
	210	0.019	0	100%				
25	240	0.027	0.002	931				
ł	270	0.04	0.006	85%				
	300	0.058	0.014	76%				
	320	0.075	0.023	694				
ļ	340	0.089	0.031	65%				
30	355	0.103	0.04	61%				

	Staphylococcus aureus 29213							
	Time T=0	Control 0	114 % Inhib					
		0_	0					
5	60		-0.003					
	120	0.003	-0.003 200%					
	165	0.006	-0.002 1334					
	210	0.01	0.002 80%					
	240	0.014	0.005 64%					
	270	0.024	0.012 50%					
10	300	0.034	0.019 44%					
	1	1						

0.06

0.072

0.09

0.102

360

380

400

420

Table 9G.

	Table 10							
20	Restoration of Ampicillin Sensitivity in an Ampicillin Resistant Strain of Escherichia coli Y1088							
20	Time T=0	Control +50 µg/ml amp	NBT 14 -	50µg/ml 1Inhib	Control -250µg/ml amp	MBT 14 amp	•250µg/ml	
	0 ·	0		0	0		0	
	60	0		o	0		0	
	120	0		0	0		0	
25	180	0		0	0		0	
	245	0		0	0.002		0	
	270	<u> </u>		. 0	0.004	0.001	75%	
	290	0.001	0.001		0.006	0.002	671	
	310	0.006	0.002	671	0.007	0.002	711	
	330	0.007	0.003	571	0.013	0.004	691	
30	355	0.013	0.005	614	0.02	0.006	70%	
	370	0.017	0.007	59%	0.022	0.008	641	
	390	0.026	0.011	58%	0.03	0.013	571	
	410	0.032	0.016	50%	0.039	0.018	544	
	430	0.038	0.021	45%	0.043	0.023	461	
35	450	0.052	0.026	50%	0.062	0.031	50%	
	470	0.069	0.035	49%	0.075	0.041	451	

0.031

0.039

0.047

0.058

0.063

35%

35%

36%

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Table 11
Number of Bacteria in the blood

	T=0	T=24 hr.	Change over 24 hours
Saline Control	1x10° bactería	3x10° bacteria	3 fold increase in bacteria
+Oligo NBT 132	1x10° bacteria	0.13x10° bacteria	10 fold reduction in bacteria

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What is claimed is:

A method for treating an animal, including a human, having an infection caused by a pathogenic bacterium, comprising: administering to the animal a composition
 comprising a pharmaceutically acceptable carrier and a substantially nuclease resistant oligonucleotide having about 8 to about 80 nucleotides and targeted to a nucleic acid or protein in the bacterium in an amount sufficient to alleviate a symptom of the infection.

- 2. The method of claim 1, wherein the nucleic acid or protein is involved in the synthesis, metabolism, assembly or regulation of at least one of the group consisting of energy, DNA replication, cell division, regulatory proteins, cell walls, sugars, virulence, fatty acids, mRNAs, tRNAs, rRNAs, ribosomal proteins, proteins involved in protein synthesis, phospholipids, periplasmic proteins, secretory proteins,
- flagellar proteins, transport proteins, amino acids, lipopolysaccharides, purines, pyrimidines, pili, outer membrane proteins, nitrogen, antibiotic binding proteins and vitamins.
- The method of claim 1, wherein the oligonucleotide is capable of associating with a nucleic acid or protein in the bacterium such that it inhibits at least one of the group consisting of bacterial growth, reproduction, metabolism,
 synthesis of toxins, progress of infection and virulence.
- 4. The method of claim 3, wherein the associating is hybridizing to an mRNA in the bacterium at or near the initiation codon, in the 5' untranslated region, in the 3' untranslated region, internal to the coding region or an 30 intermediate region of the mRNA.
 - 5. The method of claim 3, wherein the associating is hybridizing to DNA in the bacterium.
 - 6. The method of claim 5, wherein the hybridizing forms a triplex structure.
- 35 7. The method of claim 3, wherein the associating is binding with a protein in the bacterium.

8. The method of claim 1, wherein the oligonucleotide hybridizes to any one of the operons listed in Table 1.

- 9. The method of claim 1, wherein the oligonucleotide hybridizes to any one of the genes listed in Table 1.
- 5 10. The method of claim 1, wherein the oligonucleotide comprises a sequence drawn from SEQ ID NOS. 1-176 of the Sequence Listing or a functional equivalent thereof.
- 11. The method of claim 1, wherein the oligonucleotide has been purified by a method comprising at least one method 10 from the group consisting of diafiltration, gel filtration, high performance liquid chromatography, fast performance liquid chromatography, alcohol precipitations, or alcohol extractions followed by ethanol or chloroform extractions.
- 12. The method of claim 1, wherein the oligonucleotide 15 was purified by gel filtration.
 - 13. The method of claim 1, wherein the oligonucleotide is capable of inhibiting growth of the bacterium in an MIC assay.
- 14. The method of claim 1, wherein the oligonucleotide 20 has been modified in at least one base, sugar or internucleotide linkage so as to increase nuclease resistance, stability, specificity or uptake by bacteria of the oligonucleotide.
- 15. The method of claim 1, wherein the oligonucleotide 25 is selected from at least one of the group consisting of:
 - a) partially or fully substituted phosphorothicate oligonucleotides or analogues thereof;
 - b) partially or fully substituted alkyl phosphonate oligonucleotides or analogues thereof;
 - c) partially or fully substituted phosphate ester oligonucleotides or analogues thereof;

30

- d) partially or fully substituted phosphoramidate oligonucleotides or analogues thereof;
- e) partially or fully substituted 2' modified RNA
 35 oligonucleotides or analogues thereof;
 - f) partially or fully substituted morpholino oligonucleotides or analogues thereof;

- 133 -

g) partially or fully substituted peptide nucleic acid oligonucleotides or analogues thereof;

- h) partially or fully substituted dithioate oligonucleotides or analogues thereof;
- i) partially or fully substituted 5' thio oligonucleotides or analogues thereof;

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20.

- j) partially or fully substituted propyne oligonucleotides or analogues thereof;
 - k) chimerics of any combination of the above; and
- 1) any chemical modifications of the oligonucleotide which leave the oligonucleotide capable of specifically binding the nucleic acid or protein.
- 16. The method of claim 1, wherein the administration is selected from the group consisting of oral, intravenous,15 intramuscular, intraperitoneal, subcutaneous, intradermal, inhalation and topical administration.
 - 17. The method of claim 1, wherein the bacterium is gram positive.
- 18. The method of claim 1, wherein the bacterium is 20 gram negative.
 - 19. The method of claim 1, wherein the bacterium is acid fast.

The method of claim 1, wherein the bacterium is a

- member of a genus selected from the group consisting of
 25 Aerococcus, Listeria, Streptomyces, Actinomadura,
 Lactobacillus, Eubacterium, Arachnia, Mycobacterium,
 Peptostreptococcus, Staphylococcus, Corynebacterium,
 Erysipelothrix, Dermatophilus, Rhodococcus, Bifodobacterium,
- 30 Micrococcus, Kurthia, Nocardia, Nocardiopsis, Rothia, Propionibacterium, Actinomyces, Enterococcus, Pneumococcus, and Clostridia.

Lactobacillus, Streptococcus, Bacillus, Peptococcus,

- 21. The method of claim 1, wherein the bacterium is a member of the genus Staphylococcus.
- 35 22. The method of claim 21, wherein the bacterium is Staphylococcus aureus.

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23. The method of claim 1, wherein the bacterium is a member of the genus *Pseudomonas*.

- 24. The method of claim 1, wherein the bacterium is a member of the genus *Klebsiella*.
- 5 25. The method of claim 1, wherein the bacterium is a member of the genus Yersinia.
 - 26. The method of claim 1, wherein the bacterium is a member of the genus Neisseria.
- 27. The method of claim 1, wherein the bacterium is a 10 member of the genus Serratia.
 - 28. The method of claim 1, wherein the bacterium is a member of the genus Streptococcus.
 - 29. The method of claim 28, wherein the bacterium is Streptococcus pyogenes.
- 30. The method of claim 28, wherein the bacterium is Streptococcus pneumoniae.
 - 31. The method of claim 1, wherein the bacterium is a member of the genus Shigella.
- 32. The method of claim 1, wherein the bacterium is a 20 member of the genus *Haemophilus*.
 - 33. The method of claim 1, wherein the bacterium is a member of the genus Mycobacterium.
 - 34. The method of claim 1, wherein the bacterium is a member of the genus *Helicobacter*.
- 25 35. The method of claim 1, wherein the bacterium is a member of the genus *Enterococcus*.
 - 36. The method of claim 1, wherein the bacterium is a member of the genus Vibrio.
- 37. The method of claim 1, wherein the bacterium is a 30 member of the genus Salmonella.
 - 38. The method of claim 1, wherein the bacterium is a Pneumococcus.
 - 39. The method of claim 1, wherein the bacterium is Escherichia coli.
- 35 40. A composition comprising a pharmaceutically acceptable carrier and a substantially nuclease resistant oligonucleotide having about 8 to about 80 nucleotides and

targeted to a nucleic acid or protein in the bacterium in an amount sufficient to alleviate a symptom of the infection.

- 41. The composition of claim 40, wherein the nucleic acid or protein is involved in the synthesis, metabolism, 5 assembly or regulation of at least one of the group consisting of energy, DNA replication, cell division, regulatory proteins, cell walls, sugars, virulence, fatty acids, mRNAs, tRNAs, rRNAs, ribosomal proteins, proteins involved in protein synthesis, phospholipids, periplasmic 10 proteins, secretory proteins, flagellar proteins, transport proteins, amino acids, lipopolysaccharides, purines, pyrimidines, pili, outer membrane proteins, nitrogen, antibiotic binding proteins and vitamins.
- 42. The composition of claim 40, wherein the
 15 oligonucleotide is capable of associating with a nucleic acid
 or protein in the bacterium such that it inhibits at least
 one of the group consisting of bacterial growth,
 reproduction, metabolism, synthesis of toxins, progress of
 infection and virulence.
- 43. The composition of claim 42, wherein the associating is hybridizing to an mRNA in the bacterium at or near the initiation codon, in the 5' untranslated region, in the 3' untranslated region, internal to the coding region or an intermediate region of the mRNA.
- 25 44. The composition of claim 42, wherein the associating is hybridizing to DNA in the bacterium.
 - 45. The composition of claim 44, wherein the hybridizing forms a triplex structure.
- 46. The composition of claim 42, wherein the 30 associating is binding with a protein in the bacterium.
 - 47. The composition of claim 40, wherein the oligonucleotide hybridizes to any one of the operons listed in Table 1.
- 48. The composition of claim 40, wherein the 35 oligonucleotide hybridizes to any one of the genes listed in Table 1.

49. The composition of claim 40, wherein the oligonucleotide comprises a sequence drawn from SEQ ID NOS. 1-176 of the Sequence Listing or a functional equivalent thereof.

- 5 50. The composition of claim 40, wherein the oligonucleotide has been purified by a method comprising at least one method from the group consisting of diafiltration, gel filtration, high performance liquid chromatography, fast performance liquid chromatography, alcohol precipitations or alcohol extractions followed by ethanol or chloroform extractions.
 - 51. The composition of claim 40, wherein the oligonucleotide was purified by gel filtration.
- 52. The composition of claim 40, wherein the 15 oligonucleotide is capable of inhibiting growth of the bacterium in an MIC assay.
 - 53. The composition of claim 40, wherein the oligonucleotide has been modified in at least one base, sugar or internucleotide linkage so as to increase nuclease
- 20 resistance, stability, specificity or uptake by bacteria of the oligonucleotide.
 - 54. The composition of claim 40, wherein the oligonucleotide is selected from at least one of the group consisting of:
- a) partially or fully substituted phosphorothicate oligonucleotides or analogues thereof;
 - b) partially or fully substituted alkyl phosphonate oligonucleotides or analogues thereof;
- c) partially or fully substituted phosphate ester30 oligonucleotides or analogues thereof;
 - d) partially or fully substituted phosphoramidate oligonucleotides or analogues thereof;
 - e) partially or fully substituted 2' modified RNA oligonucleotides or analogues thereof;
- f) partially or fully substituted morpholino oligonucleotides or analogues thereof:

g) partially or fully substituted peptide nucleic acid oligonucleotides or analogues thereof;

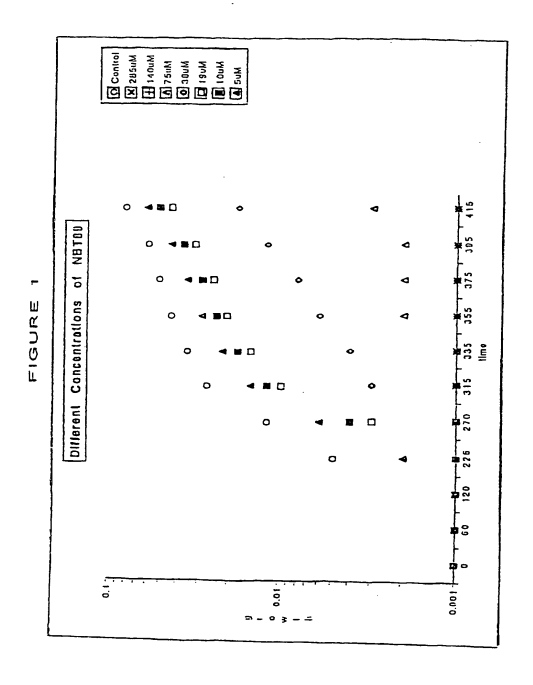
- h) partially or fully substituted dithioate oligonucleotides or analogues thereof;
- 5 i) partially or fully substituted 5' thio oligonucleotides or analogues thereof;
 - j) partially or fully substituted propyne oligonucleotides or analogues thereof;
 - k) chimerics of any combination of the above; and
- 10 l) any chemical modifications of the oligonucleotide which leave the oligonucleotide capable of specifically binding the nucleic acid or protein.
 - 55. The composition of claim 40, wherein the bacterium is gram positive.
- 15 56. The composition of claim 40, wherein the bacterium is gram negative.
 - 57. The composition of claim 40, wherein the bacterium is acid fast.
- 58. The composition of claim 40, wherein the bacterium 20 is a member of a genus selected from the group consisting of Aerococcus, Listeria, Streptomyces, Actinomadura, Lactobacillus, Eubacterium, Arachnia, Mycobacterium, Peptostreptococcus, Staphylococcus, Corynebacterium, Erysipelothrix, Dermatophilus, Rhodococcus, Bifodobacterium,
- 25 Lactobacillus, Streptococcus, Bacillus, Peptococcus,
 Micrococcus, Kurthia, Nocardia, Nocardiopsis, Rothia,
 Propionibacterium, Actinomyces, Enterococcus, Pneumococcus,
 and Clostridia.
- 59. The composition of claim 40, wherein the bacterium 30 is a member of the genus Staphylococcus.
 - 60. The composition of claim 40, wherein the bacterium is Staphylococcus aureus.
 - 61. The composition of claim 40, wherein the bacterium is a member of the genus *Pseudomonas*.
- 35 62. The composition of claim 40, wherein the bacterium is a member of the genus *Klebsiella*.

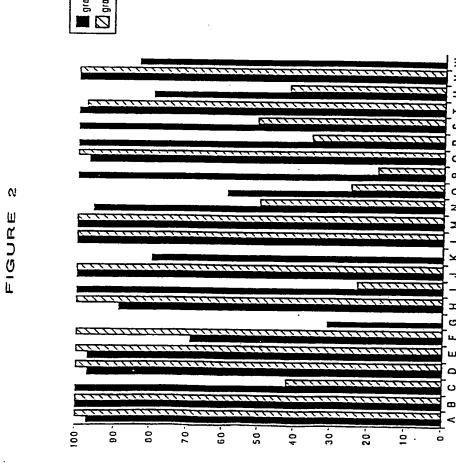
63. The composition of claim 40, wherein the bacterium is a member of the genus *Yersinia*.

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- 64. The composition of claim 40, wherein the bacterium is a member of the genus *Neisseria*.
- 5 65. The composition of claim 40, wherein the bacterium is a member of the genus Serratia.
 - 66. The composition of claim 40, wherein the bacterium is a member of the genus Streptococcus.
- 67. The composition of claim 66, wherein the bacterium 10 is Streptococcus pyogenes.
 - 68. The composition of claim 66, wherein the bacterium is Streptococcus pneumoniae.
 - 69. The composition of claim 40, wherein the bacterium is a member of the genus Shigella.
- 70. The composition of claim 40, wherein the bacterium is a member of the genus *Haemophilus*.
 - 71. The composition of claim 40, wherein the bacterium is a member of the genus Mycobacterium.
- 72. The composition of claim 40, wherein the bacterium 20 is a member of the genus *Helicobacter*.
 - 73. The composition of claim 40, wherein the bacterium is a member of the genus *Enterococcus*.
 - 74. The composition of claim 40, wherein the bacterium is a member of the genus *Vibrio*.
- 75. The composition of claim 40, wherein the bacterium is a member of the genus Salmonella.
 - 76. The composition of claim 40, wherein the bacterium is Escherichia coli.
- 77. The composition of claim 40, wherein the bacterium 30 is *Pneumococcus*.
 - 78. A compound, comprising:
 - a) an antibiotic; and
- b) a substantially nuclease resistant oligonucleotide having about 8 to about 80 nucleotides and targeted to a
 35 nucleic acid or protein in a bacterium,
 - wherein said antibiotic is covalently linked to said oligonucleotide.

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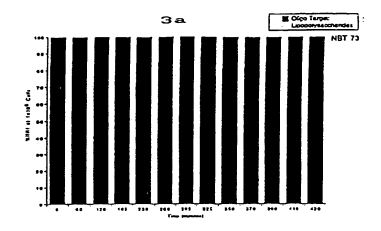


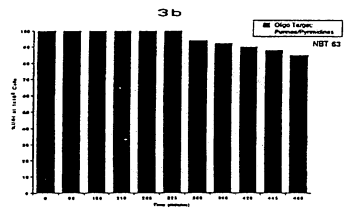


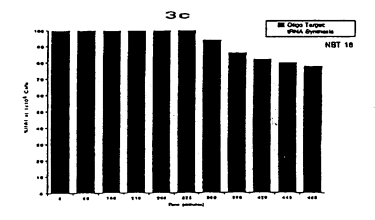
%INH at 1x108 Cells

3 / 2 0 Growth Inhibition of Bacterial Strains with Oligos



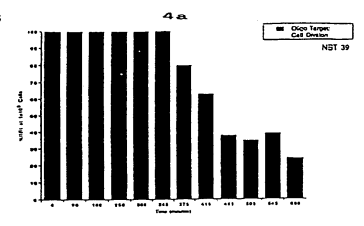


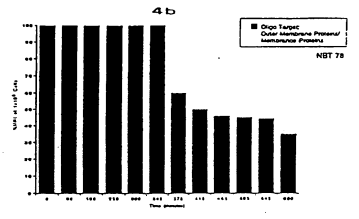


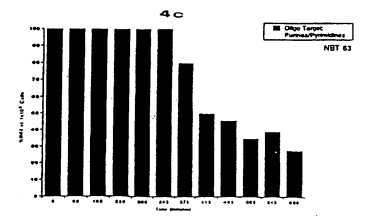


 $\mbox{4/20}$ Growth Inhibition of Bacterial Strains with Oligos

Pseudomonas



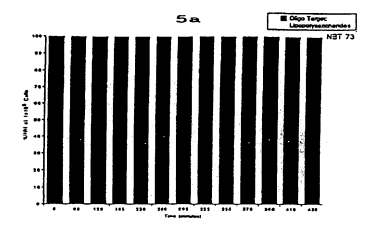


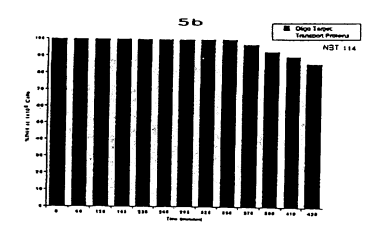


5 / 2 0

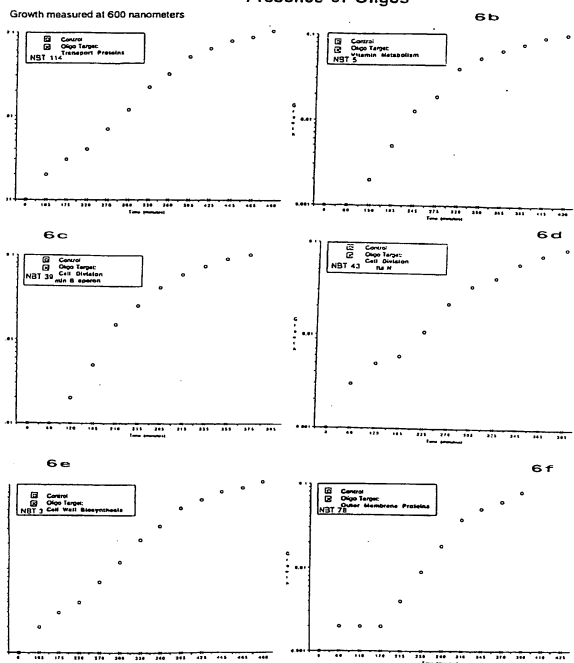
Growth Inhibition of Bacterial Strains with Oligos

Klebsiella





Growt. of E. coli 35218 (multiple drug ...sistance) in the Presence of Oligos

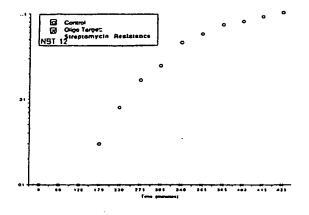


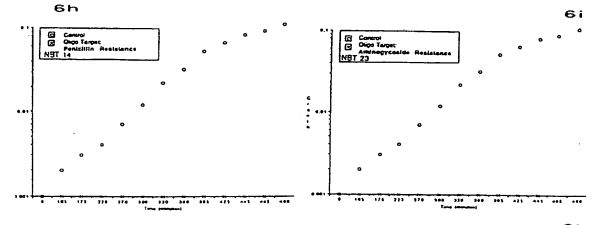
WO 98/03533 7 / 2 0 PCT/US97/12961

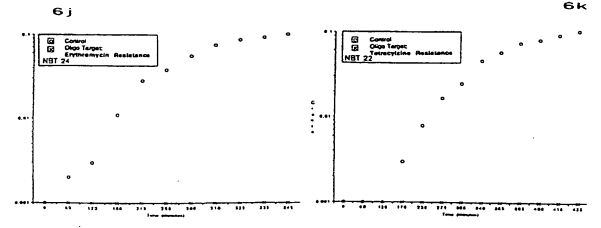
Growth of E. coli 35218 (multiple drug resistance) in the Presence of Oligos

Growth measured at 600 nanometers

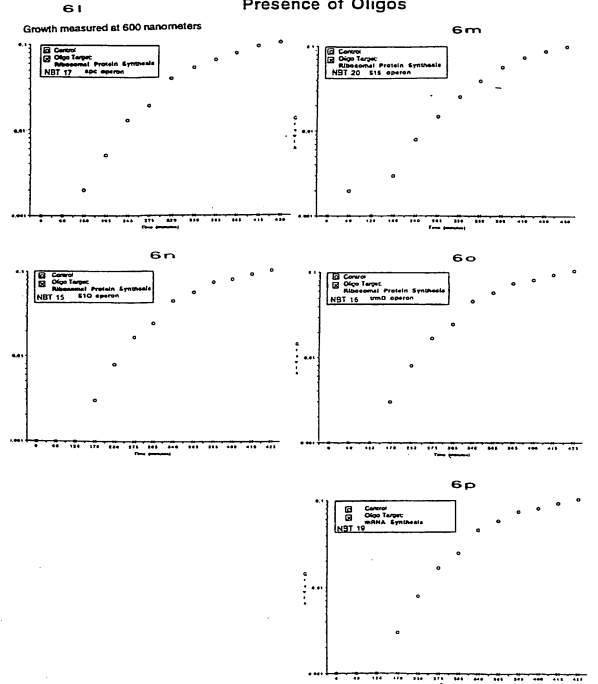
6g



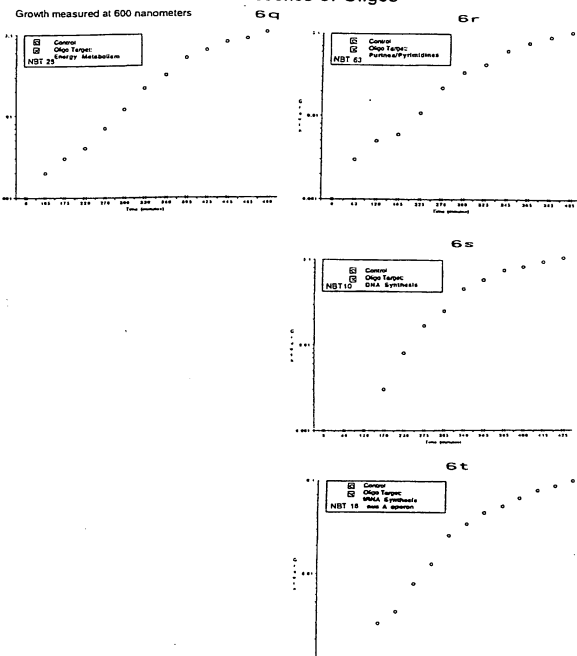




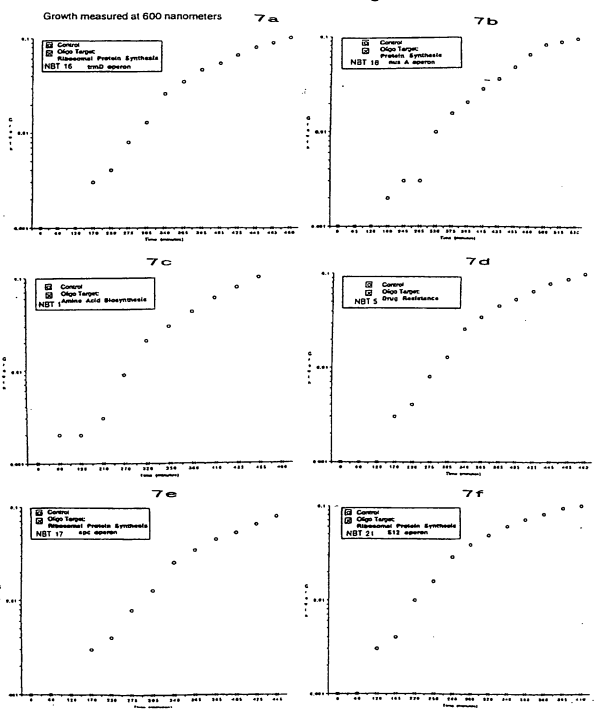
Growth of E. coli 35218 (multiple drug resistance) in the Presence of Oligos



Growth of E. coli 35218 (multiple drug resistance) in the Presence of Oligos

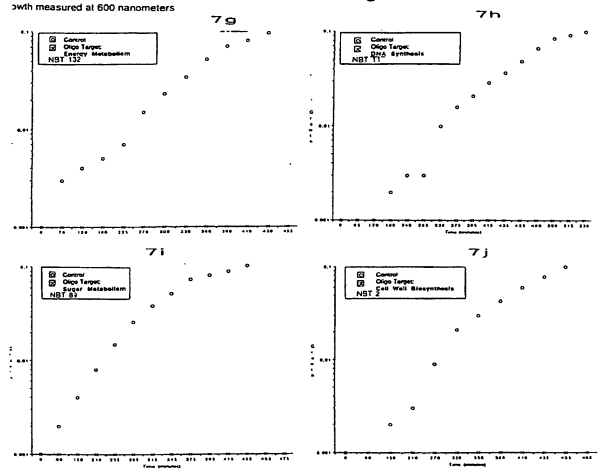


Growth In. libition of Staph 13301 (penicular resistant) in the Presence of Oligos



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Growth Inhibition of Staph 13301 (penicillin resistant) in the Presence of Oligos



12/20 Animal Data

A) Lister Model

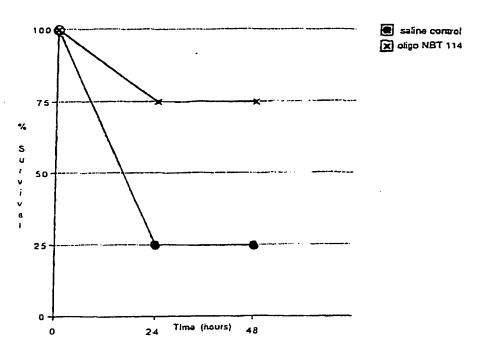


FIGURE 8

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In Vivo Efficacy

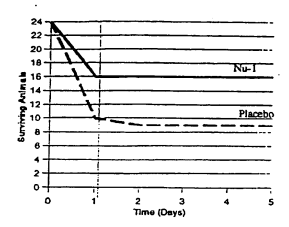


FIGURE 9

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Standard Overnight MIC Assay- Staph. aureus 3 Day Time Course

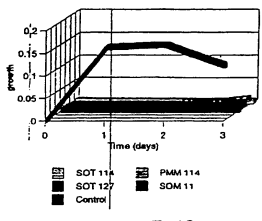
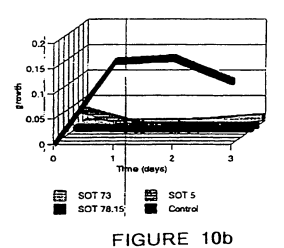


FIGURE 10a



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Standard Overnight MIC Assay Serratia liquefaciens

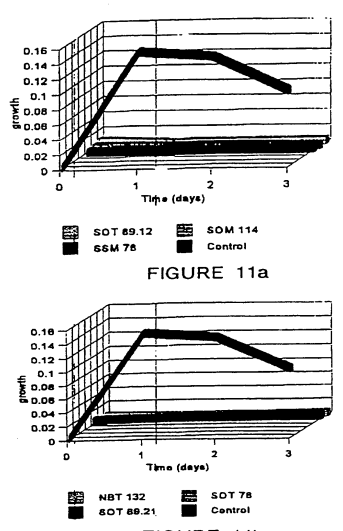
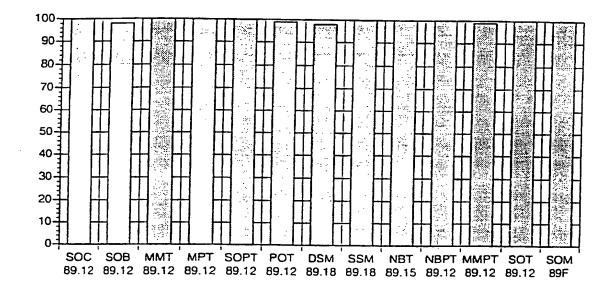


FIGURE 11b

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FIGURE 12

Standard MIC Assay Staph. aureus



SOC - 5' - 6 Ds 6Mo - Cholesteryl - 3'

SOB - 5' - 6 Ds 6Mo - Biotin - 3'

MMT - 5' - 12 Mo Invert T - 3'

MPT - 5' - 10 Mo 2Mp Invert T - 3'

SOPT - 5' - 6 Ds 4 Mo 2Mp Invert T - 3'

POT -5' - 12 Po (Invert T) - 3'

DSM - 5' - 8 Ds 10 Ms 1 Do - 3'

SSM - 5' - 18 Ms 1 Do - 3'

NBT - 5' - 14 Ds Do - 3'

NBPT - 5' - 10 Ds 2Mp Invert T - 3'

MMPT - 5' - 10 Mo 2 Mp Invert T - 3'

SOT - 5' - 6 Ds 6Mo Invert T - 3'

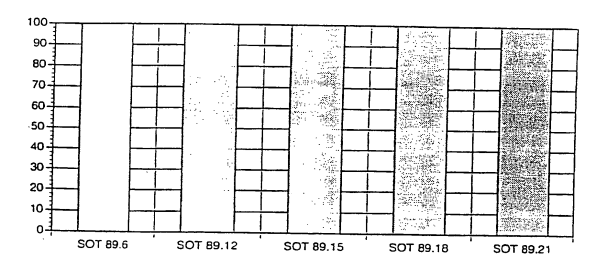
SOM-F - 5' - 1 Ms 4Ds 12 Mo 3 Ms 1 Do - 3'

Different constructs that work well in inhibition of bacterial growth.

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FIGURE 13

Standard MIC Assay Staph. aureus

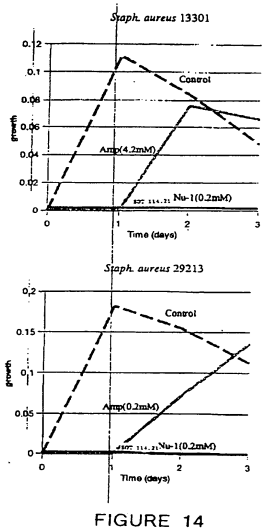


SOT 89.6-6mer SOT 89.12-12 mer SOT 89.15-15 mer SOT 89.18-18 mer SOT 89.21-21 mer

Oligos of different lengths work well in inhibition of bacterial growth.

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Comparison of Oligo 114 and Ampicillin



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Pseudomonas aeroginosa 10145

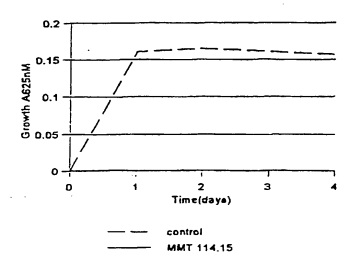


FIGURE 15

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Streptococcus pyogenes 14289

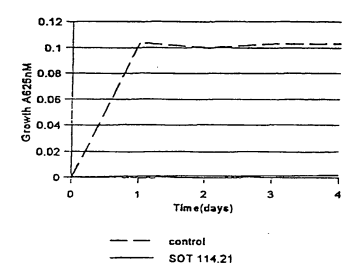


FIGURE 16

INTERNATIONAL SEARCH REPORT

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A CLASS	MFICATION OF SUBJECT MATTER C07H21/00 A61K31/70 C12N	15/11	C	
According	to International Pelent Classification (IPC) or to both national cla	seification and IPC	·	
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Minimum d IPC 6	tocumentation searched (classification system followed by classi CO7H A61K C12N	fication symbols)		
Documents	ation searched other than minimum documentation to the extent t	hat such documents are included in the fielde se	arched	
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C. DOCUM	ENTS CONSIDERED TO BE RELEVANT			
Gategory*	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.	
X	US 5 294 533 A (LUPSKI JAMES R March 1994 cited in the application see the whole document	ET AL) 15	1-77	
P,X	WO 96 29399 A (SOD CONSEILS RE;PIROTZKY EDUARDO (FR); COLOTE 26 September 1996 see the whole document	1-77		
A	L. A. CHRISEY ET AL.: "Intern Oligodeoxyribonucleotides by V parahaemolyticus" ANTISENSE RES. DEV., vol. 3, 1993, pages 367-381, XP002045887 cited in the application			
		-/		
X Furt	her documents are listed in the continuation of box C.	X Patent family members are listed in	n annex.	
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	UON) DOCUMENTS CONSIDERED TO BE RELEVANT		
lagory *	Castion of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
	M.A. RAHMAN ET AL.: "Antibacterial Activity and Inhibition of Protein Synthesis in Escherichia coli by Antisense DNA Analogs" ANTISENSE RES. DEV., vol. 1, 1991, pages 319-27, XP002045888 cited in the application		
	K. JAYARAMAN ET AL.: "Selective Inhibition of Escherichia coli Protein Synthesis and Growth by Nonionic Oligonucleotides Complementary to the 3' End of 16S rRNA" PROC. NATL. ACAD. SCI. USA, vol. 78, 1981, pages 1537-41, XP002045889 cited in the application		
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INTERNATIONAL SEARCH REPORT

Information on patent family members

Internat. ..al Application No PCT/US 97/12961

Patent document cited in search report	Publication date	Patent (amily member(s)	Publication date '
US 5294533 A	15-03-94	AU 645339 B	13-01-94
		AU 8263491 A	27-02-92
	•	CA 2048450 A	24-02-92
		EP 0472434 A	26-02-92
		JP 6303977 A	01-11-94
		AT 137806 T	15 - 05-96
		AU 4180889 A	05-02-90
		DE 68926455 D	13-06-96
		DE 68926455 T	31-10-96
		EP 0424473 A	02-05-91
		JP 3505672 T	12-12-91
		WO 9000624 A	25-01-90
WO 9629399 A	26-09-96	AU 5149796 A	08-10-96

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C07H 21/00, A61K 31/70, C12N 15/11	A1	(43) International Publication Date: 29 January 1998 (29.01.98)
 21) International Application Number: PCT/US 22) International Filing Date: 23 July 1997 (30) Priority Data: 08/685,575 24 July 1996 (24.07.96) 71) Applicant: OLIGOS ETC. AND OLIGOS THERAP INC. (US/US); 29970 S.W. Town Center L Wilsonville, OR 97070 (US). 72) Inventors: ARROW, Amy; 15 Equestrian Ridge Roton, CT 06470 (US). DALE, Roderic, M., K.; 26' 45th Drive, Wilsonville, OR 97070 (US). THO Theresa, L.; 2222 S.W. Ek Road, West Linn, O (US). 74) Agents: FRIEBEL, Thomas, E. et al.; Pennie & Edmo 1155 Avenue of the Americas, New York, NY 100 	(23.07.9 PEUTIC Loop W. ad, Nev 761 S.V MPSON R 9706	CA, CN, CU, CZ, EE, GE, GH, HU, IL, IS, JP, KG, KP, KR, KZ, LC, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UZ, VN, YU, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG). Published With international search report. With amended claims. Date of publication of the amended claims: 2 April 1998 (02.04.98)

(57) Abstract

A novel method is provided that teaches the therapeutic use of nuclease resistant oligonucleotides for treating animals having an infection caused by a pathogenic bacterium. The method involves the integration of (1) methods for selecting the correct oligonucleotide, (2) synthesis and purification of nuclease resistant oligonucleotides, and (3) methods for *in vitro* analysis of potential antimicrobial oligonucleotides. The described oligonucleotides may comprise modified backbones, sugar residues, bases, or mixtures and have been subject to purification resulting in oligonucleotides that are capable of inhibiting the growth of a broad spectrum of clinically relevant bacterial species.

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AMENDED CLAIMS

[received by the International Bureau on 12 February 1998 (12.02.98); original claims 1-78 replaced by new claims 1-21 (2 pages)]

- 1. The use of a substantially nuclease resistant oligonucleotide having about 8 to about 80 nucleotides in the preparation of a medication for the treatment of infection by pathogenic bacteria.
- 2. The use of claim 1 wherein said bacteria are gram positive.
- 3. The use of claim 2 wherein said bacteria is selected from the group consisting of: Aerococcus, Listeria, Streptomyces, Actinomadura, Lactobacillus, Eubacterium, Arachnia, Mycobacterium, Peptostreptococcus, Corynebacterium, Erysipelothrix, Dermatophilus, Rhodococcus, Bifodobacterium, Lactobacillus, Bacillus, Peptococcus, Micrococcus, Kurthia, Nocardia, Nocardiopsis, Rothia, Propionibacterium, Actinomyces, Pneumococcus, and Clostridia.
- 4. The use of claim 2, wherein the bacterium is a member of the genus Staphylococcus.
- 5. The use of claim 4, wherein the bacterium is Staphylococcus aureus.
- 6. The use of claim 2, wherein the bacterium is a member of the genus Streptococcus.
- 7. The use of claim 6, wherein the bacterium is Streptococcus pyogenes.
- 8. The use of claim 6, wherein the bacterium is Streptococcus pneumoniae.
- 9. The use of claim 2, wherein the bacterium is a member of the genus *Enterococcus*.

1 4 O AMENDED SHEET (ARTICLE 19)

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10. The use of claim 1 wherein said bacteria are gram negative.

- 11. The use of claim 10, wherein the bacterium is a member of the genus *Pseudomonas*.
- 12. The use of claim 10, wherein the bacterium is a member of the genus *Klebsiella*.
- 13. The use of claim 10, wherein the bacterium is a member of the genus Yersinia.
- 14. The use of claim 10, wherein the bacterium is a member of the genus Neisseria.
- 15. The use of claim 10, wherein the bacterium is a member of the genus Serratia.
- 16. The use of claim 10, wherein the bacterium is a member of the genus Shigella.
- 17. The use of claim 10, wherein the bacterium is a member of the genus Haemophilus.
- 18. The use of claim 10, wherein the bacterium is a member of the genus Mycobacterium.
- 19. The use of claim 10, wherein the bacterium is a member of the genus *Vibrio*.
- 20. The use of claim 10, wherein the bacterium is a member of the genus Salmonella.
- 21. The use of claim 10, wherein the bacterium is Escherichia coli.

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